

STIC Search Report Biotech-Chem Library

STIC Database Tracking Number: 203825

TO: Rip A Lee

Location: Remsen 10a24

Wednesday, October 04, 2006

Art Unit: 1713

Phone: 571-272-1104

Serial Number: 10 / 541644

From: Jan Delaval Location: EIC 1700

Remsen 4a30

Phone: 571-272-2504

jan.delaval@uspto.gov

Search Notes



OCT 4 RECD SEARCH REQUEST FORM

Pat. & T.M. $Oti_{C\Theta}$ Scientific and Technical Information Center.

Requester's Full Name: LET, LPA. Examiner #: 7560 Date: Octobr 04, 2006 Art Unit: 1710 Phone Number 39 2-1104 Serial Number: 10, 541, 644 Mail Box and Bldg/Room Location: DEM 10 A24 Results Format Preferred (circle): PAPER DISK E-MAIL
If more than one search is submitted, please prioritize searches in order of need.
Please provide a detailed statement of the search topic, and describe as specifically as possible the subject matter to be searched. Include the elected species or structures, keywords, synonyms, acronyms, and registry numbers, and combine with the concept or attility of the invention. Define any terms that may have a special meaning. Give examples or relevant citations, authors, etc, if known. Please attach a copy of the cover sheet, pertinent claims, and abstract.
Title of Invention: GRUP 3 BEDGED METALLO COVE
inventors (please provide full names): <u>CARPENTIER</u> , J.F. REZAVI, A. KIPILLOV, E.
Earliest Priority Filing Date:
For Sequence Searches Only Please include all pertinent information (parent, child, divisional, or issued patent numbers) along with the ppropriate serial number.
Please scorch for yttrium or scandium metallocenes with following structure
cyclopentedienyl
P Y, Sc Hurrenyl
Dis bridging vait Messi, Mesch citz.
TAFF USE ONLY Type of Search Vendors and cost where applicable
NA Sequence (#) STN
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e Searcher Picked Up: 1014 0
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ical Prep Time: Patent Family WWW/Internet
ine Time: Cuber (Creation)

=> fil reg FILE 'REGISTRY' ENTERED AT 15:15:03 ON 04 OCT 2006 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2006 American Chemical Society (ACS)

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STRUCTURE FILE UPDATES: 3 OCT 2006 HIGHEST RN 909488-17-1 DICTIONARY FILE UPDATES: 3 OCT 2006 HIGHEST RN 909488-17-1

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH June 30, 2006

Please note that search-term pricing does apply when conducting SmartSELECT searches.

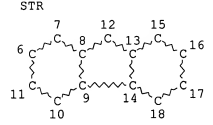
REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

=> d sta que 132

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L26 1459387 SEA FILE=REGISTRY ABB=ON PLU=ON L25 OR ((Y OR SC OR LA OR ND
OR SM)/ELS OR (?YTTRIUM? OR ?LANTHANUM? OR ?NEODYMIUM? OR
?SAMARIUM? OR ?SCANDIUM?)/CNS)

L27



NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 18

STEREO ATTRIBUTES: NONE

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L30 STR

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STEREO ATTRIBUTES: NONE

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919 ANSWERS

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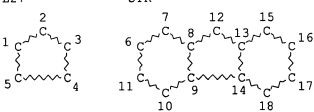
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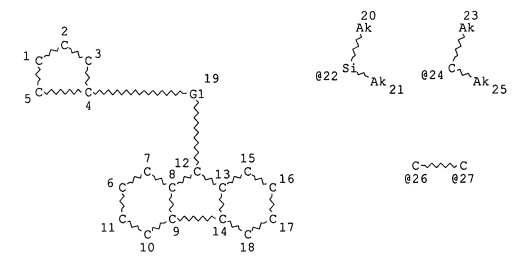
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L48 STR



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DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

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STEREO ATTRIBUTES: NONE

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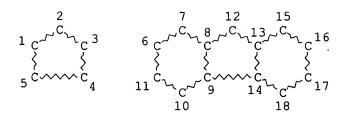
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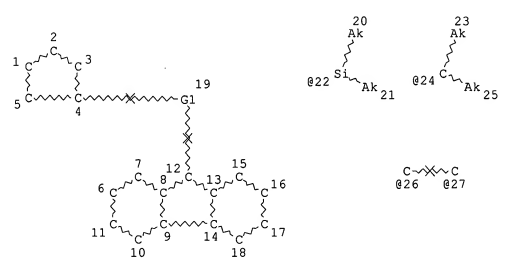
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CONNECT IS E1 RC AT 23

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DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

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NUMBER OF NODES IS 27

STEREO ATTRIBUTES: NONE

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100.0% PROCESSED 2793 ITERATIONS

SEARCH TIME: 00.00.01

947 ANSWERS

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L20

L36

L37

L38

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                E KIRILLOV/AU
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L4
            152 S E3-E23
                E RAZAVI/AU
L5
            130 S E4-E7
                E FINA/PA,CS
L6
           1045 S E3, E4
                E ATOFINA/PA, CS
L7
            845 S E3, E4
                E GROUP III/CW, CT
\Gamma8
         976263 S E40+OLD, NT OR E45+OLD, NT
L9
         556594 S E128+OLD, NT
L10
         264688 S E144+OLD, NT OR E142+OLD, NT
L11
          18751 S (GROUP IIIA? OR GROUP IIIB?)/CT
L12
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L13
             54 S L12 AND ?CYCLOPENT? AND ?FLUOREN?
L14
            130 S L12 AND ?METALLOCEN?
L15
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L16
L17
             13 S L16 AND (?YTTRIUM? OR ?LANTHANUM? OR ?NEODYMIUM? OR ?SAMARIUM
L18
             14 S L15, L17
L19
             14 S L1, L18
                SEL RN
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FILE 'REGISTRY' ENTERED AT 14:37:32 ON 04 OCT 2006

162 S E1-E162

11 S L32 AND ND/ELS

17 S L32 AND SM/ELS

119 S L33-L37 NOT (AYS OR TIS)/CI

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L21
             62 S L20 AND CCS/CI
L22
             45 S L21 AND (?YTTRIUM? OR ?LANTHANUM? OR ?NEODYMIUM? OR ?SAMARIUM
L23
             45 S L21 AND (Y OR SC OR LA OR ND OR SM)/ELS
L24
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L25
        1402823 S E6 OR E15 OR E25
L26
        1459387 S L25 OR ((Y OR SC OR LA OR ND OR SM)/ELS OR (?YTTRIUM? OR ?LAN
L27
L28
             50 S L27 SAM SUB=L26
L29
           2794 S L27 FUL SUB=L26
                SAV L29 LEE541/A
L30
                STR L27
L31
             50 S L30 SAM SUB=L29
L32
            919 S L30 FUL SUB=L29
                SAV L32 LEE541A/A
L33
             80 S L32 AND Y/ELS
L34
             77 S L32 AND SC/ELS
L35
             63 S L32 AND LA/ELS
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L39
             58 S L38 NOT ?FULLER?/CNS
L40
             56 S L39 NOT (C82 OR C80)
L41
             34 S L20 AND L29
L42
             17 S L41 AND (Y OR SC)/ELS
                SEL RN 2 3 6 7 11 12
L43
             11 S L42 NOT E1-E6
L44
             30 S L40 AND (Y OR SC)/ELS
L45
             19 S L44 NOT L42
                SEL RN 9 11-15
             13 S L45 NOT E7-E12
T.46
1.47
             24 S L43, L46
L48
                STR L30
L49
              1 S L48 SAM SUB=L29
L50
             29 S L48 FUL SUB=L29
                SAV L50 LEE541B/A
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L66
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             20 S L63, L67
L68
L69
              9 S L63 AND L65
L70
             16 S L67, L69
L71
              4 S L63 NOT L70
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FILE 'REGISTRY' ENTERED AT 15:15:03 ON 04 OCT 2006

=> fil hcaplus FILE 'HCAPLUS' ENTERED AT 15:15:34 ON 04 OCT 2006 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 4 Oct 2006 VOL 145 ISS 15 FILE LAST UPDATED: 3 Oct 2006 (20061003/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

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ANSWER 1 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN
ΑN
     2005:140001 HCAPLUS
DN
     142:219710
ΤI
     Catalytic system for obtaining copolymers of conjugated diene(s) and
     monoolefin(s) copolymers and these copolymers
IN
     Boisson, Christophe; Monteil, Vincent; Spitz, Roger
PΑ
     Societe de Technologie Michelin, Fr.; Michelin Recherche et Technique Sa;
     Atofina
SO
     Fr. Demande, 47 pp.
     CODEN: FRXXBL
DT
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LA
     French
FAN.CNT 1
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     PATENT NO.
                        KIND
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os
     MARPAT 142:219710
AB
     The catalytic system for the title use comprises: (i) [P(Cp)(F1)Ln(X)(Lx)]
     (I) where Ln represents a lanthanide atom to which a mol. of ligand is
     connected including groups cyclopentadienyl (Cp) and fluorenyl (F1)
     connected to each other by a bridge P of formula: MR1R2, where M is an
     element of Group IVA and where R1 and R2, identical or different,
     represent each alkyl group having from 1 to 20 atoms of carbon or
     cycloalkyl groups or aryl groups having from 6 to 20 carbon atoms, where X
     represents an atom of halogen which can be chlorine, the fluorine, bromine
     or iodine, where L includes groups such as an ether, and possibly a mol.
     appreciably less chelating, such as toluene, where p is \geq 1 and x
     ≥ 0, and (ii) a cocatalyst selected from alkylmagnesium,
```

alkyllithium, alkylaluminium, Grignard reactants or their mixts.

polymers prepared using these catalyst systems have mol. weight >30,000, diene unit content >40 mol%, and C3-18 olefin unit content ≥10%. A typical I was manufactured by reaction of Me2SiC5H5C13H9 with BuLi in THF and complexation of the intermediate with NdCl3 in THF.

ΙT 839680-86-3P

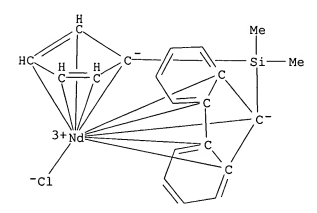
> RL: CAT (Catalyst use); IMF (Industrial manufacture); PREP (Preparation); USES (Uses)

(catalysts based on Group IVA element-bridged

fluorenylcyclopentadienyllanthanide complexes for obtaining copolymers of conjugated diene(s) and C3-18 α -olefin(s) copolymers)

RN 839680-86-3 HCAPLUS

CN Neodymium, chloro[\(\eta 10-2, 4-cyclopentadien-1-\)ylidene(dimethylsilylene)-9H-fluoren-9-ylidene]- (9CI) (CA INDEX NAME)



RETABLE

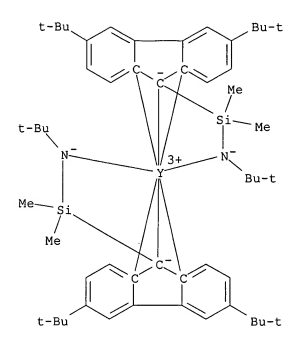
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Boisson, C Mitsui Chemicals Inc	2003 1999		US 6569799 B1 EP 0891993 A	HCAPLUS HCAPLUS
Wilson, J	[2002]	1	US 6348555 B1	HCAPLUS

- L70 ANSWER 2 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN
- AN 2004:738468 HCAPLUS
- DN 141:243979
- TΙ Catalytic components with constrained geometry comprising a fluorenyl ligand and based on Group IIIB metals
- IN Razavi, Abbas; Carpentier, Jean Francois; Kirillov, Evgueni
- PA Atofina Research, Belg.; Centre National De La Recherche Scientifique CNRS
- SO Fr. Demande, 25 pp. CODEN: FRXXBL
- DT Patent
- French LA
- FAN.CNT 1

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     WO 2004-EP2378
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     Metallocene complexes of Group IIIB metals having a fluorenyl ligand
AB
     bonded to a hydrocarbyl-substituted Si which is, in turn, bonded to a
     (substituted) N so as to constrain the geometry of the fluorenyl ligand
     are useful for catalysts in controlled polymerization of polar or nonpolar
     monomers. A typical catalyst was manufactured by stirring a suspension of 1.73
     mmol 338 mg YCl3 in THF 2 h with 5.2 mmol LiCH2SiMe3 in pentane at
     0°, removing residual LiCl, adding 1.42 mmol 3,6-di-tert-butyl-9-
     (tert-butyldimethylsilyl)fluorene in pentane at 0°, warming to room
     temperature, and stirring 30 h.
ΙT
     624739-65-7P 624739-67-9P 624739-71-5P
     752997-00-5P
     RL: CAT (Catalyst use); IMF (Industrial manufacture); PREP (Preparation);
     USES (Uses)
        (catalysts with constrained geometry comprising fluorenyl ligands and
        based on Group IIIB metals for polymerization of polar and nonpolar monomers)
     624739-65-7 HCAPLUS
RN
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CN
     3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-
     dimethylsilanaminato(2-)-kN]yttrate(1-) (9CI) (CA INDEX NAME)
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          1
     CRN
         624739-64-6
         C54 H78 N2 Si2 Y
     CMF
     CCI CCS
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CM 2

CRN 48186-27-2 CMF C16 H32 Li O4 CCI CCS

RN 624739-67-9 HCAPLUS

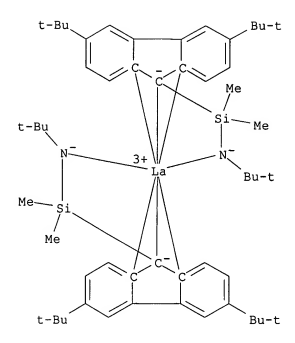
CN Lithium(1+), tetrakis(tetrahydrofuran)-, (T-4)-, bis[1-[(8a,9,9a- η)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κ N]lanthanate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 624739-66-8

CMF C54 H78 La N2 Si2

CCI CCS



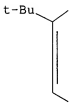
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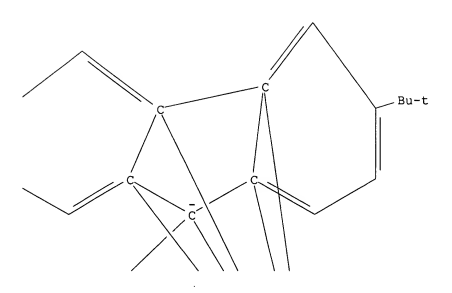
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CN Neodymium, bis[1-[(4a,4b,8a,9,9a- η)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κ N]di- μ -chlorobis(tetrahydrofuran)di-, stereoisomer (9CI) (CA INDEX NAME)

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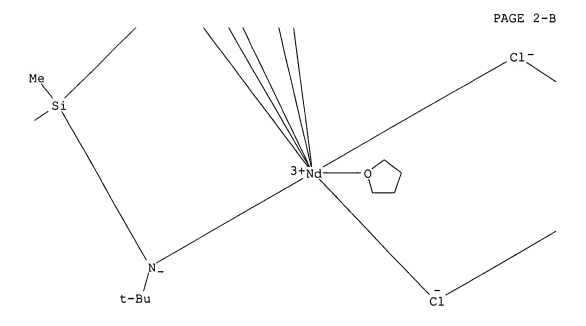


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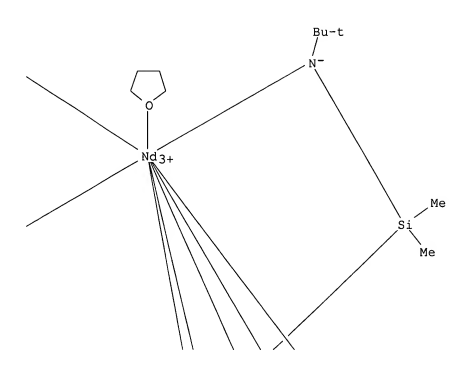


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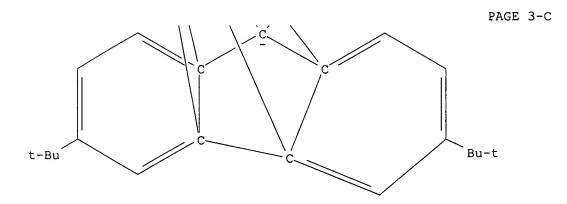
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PAGE 2-C



PAGE 3-A



RN 752997-00-5 HCAPLUS

CN Lithium(1+), bis[1,1'-oxybis[ethane]]-, bis[1-[(8a,9,9a- η)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κ N]lanthanate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 624739-66-8

CMF C54 H78 La N2 Si2

CCI CCS

2 CM

78127-97-6 CRN CMF C8 H20 Li O2 CCI CCS

L70 ANSWER 3 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

ΑN 2004:611929 HCAPLUS

DN 141:157620

Metallocenes bridged with Group III elements and based on ΤI cyclopentadienyl-fluorenyl ligands

IN Carpentier, Jean Francois; Kirillov, Evgueni;

Razavi, Abbas

Atofina Research, Belg.; Centre National de la Recherche PA Scientifique CNRS

SO Fr. Demande, 20 pp.

CODEN: FRXXBL

DT Patent

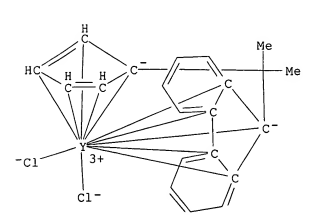
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FAN.CNT 1

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jan delaval - 4 october 2006

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OS
     MARPAT 141:157620
AB
     (FluR''Cp)M(\eta 3-C3R'5) (ether)n, in which Cp is (substituted)
     cyclopentadienyl, Flu is (substituted) fluorenyl, R'' is a structural
     bridge between Cp and Flu conferring the stereorigidity of the component,
     M is a metal of Group IIIB of the Periodic Table, each R' is identical or
     different and represents hydrogen or a hydrocarbyl comprising from 1 to 20
     atoms of carbon and n is 0, 1 or 2 are manufactured for use as catalysts for
     controlled polymerization of polar and nonpolar monomers. A typical metallocene
     was manufactured by adding 2 equiv BuLi (4.6 mL solution 1.6 M in hexane) to Et20
     containing 1 g C13H8H-CHMe2-C5H4H at -10° with vigorous stirring, aging
     the mixture 3 h at room temperature, cooling the resulting suspension to
     -20°, adding a suspension of YCl3(THF) (prepared from 0.72 g YCl3) in
     Et20, warming to room temperature, suspending 0.390 resulting powder with PhMe,
     adding 0.27 mL solution of 2 M allyl magnesium chloride in THF, and stirring
     8 h.
ΙT
     611233-16-0P
     RL: IMF (Industrial manufacture); RCT (Reactant); PREP (Preparation); RACT
     (Reactant or reagent)
        (catalyst precursor; metallocenes bridged with Group IIIB elements and
        based on cyclopentadienyl-fluorenyl ligands for catalysts for polymerization
        of polar and nonpolar monomers)
RN
     611233-16-0 HCAPLUS
CN
     Lithium(1+), [1,1'-oxybis[ethane]]tris(tetrahydrofuran)-, (T-4)-,
     dichloro[\pi10-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-
     fluoren-9-ylidene]yttrate(1-) (9CI) (CA INDEX NAME)
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611233-15-9

C21 H18 C12 Y

CRN CMF

CCI CCS

CM 2

CRN 444121-94-2 CMF C16 H34 Li O4 CCI CCS

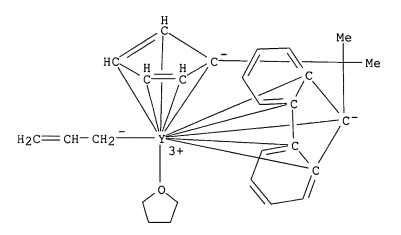
IT 714977-58-9P

RL: CAT (Catalyst use); IMF (Industrial manufacture); PREP (Preparation); USES (Uses)

(metallocenes bridged with Group IIIB elements and based on cyclopentadienyl-fluorenyl ligands for catalysts for polymerization of polar and nonpolar monomers)

RN 714977-58-9 HCAPLUS

CN Yttrium, [\eta 10-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene]-2-propenyl(tetrahydrofuran)- (9CI) (CA INDEX NAME)



RETABLE

Referenced Author | Year | VOL | PG | Referenced Work | Referenced (RAU) | (RPY) | (RVL) | (RPG) | (RWK) | File | File | Yasuda, H | 1995 | | JP 07258319 A | HCAPLUS

L70 ANSWER 4 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

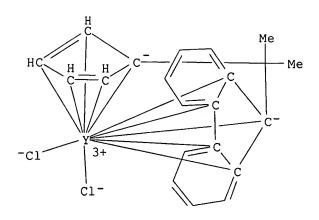
AN 2004:549514 HCAPLUS

DN 141:89539

TI Metallocenes bridged with Group III elements and based on cyclopentadienyl-fluorenyl ligands

IN Carpentier, Jean Francois; Kirillov, Evgueni;
Razavi, Abbas

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PA
    Atofina Research, Belg.; Centre National De La Recherche
     Scientifique CNRS
SO
     Fr. Demande, 22 pp.
     CODEN: FRXXBL
DТ
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LA
     French
FAN.CNT 1
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     PATENT NO.
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OS
    MARPAT 141:89539
AΒ
     (FluR''Cp)M(η3-C3R'5)(ether)n, in which Cp is (substituted)
    cyclopentadienyl, Flu is (substituted) fluorenyl, R'' is a structural
     bridge between Cp and Flu conferring the stereorigidity of the component,
    M is a metal of Group IIIB of the Periodic Table, each R' is identical or
     different and represents hydrogen or a hydrocarbyl comprising from 1 to 20
     atoms of carbon and n is 0, 1 or 2 are manufactured for use as catalysts for
     controlled polymerization of polar and nonpolar monomers. A typical metallocene
     was manufactured by adding 2 equiv BuLi (4.6 mL solution 1.6 M in hexane) to Et20
     containing 1 g C13H8H-CHMe2-C5H4H at -10° with vigorous stirring, aging
     the mixture 3 h at room temperature, cooling the resulting suspension to
     -20°, adding a suspension of YCl3(THF) (prepared from 0.72 g YCl3) in
     Et20, warming to room temperature, suspending 0.390 resulting powder with PhMe,
     adding 0.27 mL solution of 2 M allyl magnesium chloride in THF, and stirring
     8 h.
ΙT
     611233-16-0P
     RL: IMF (Industrial manufacture); RCT (Reactant); PREP (Preparation); RACT
     (Reactant or reagent)
        (catalyst precursor; metallocenes bridged with Group IIIB elements and
       based on cyclopentadienyl-fluorenyl ligands for catalysts for polymerization
       of polar and nonpolar monomers)
RN
     611233-16-0 HCAPLUS
     Lithium(1+), [1,1'-oxybis[ethane]]tris(tetrahydrofuran)-, (T-4)-,
CN
     dichloro(η10-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-
     fluoren-9-ylidene]yttrate(1-) (9CI) (CA INDEX NAME)
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         1
     CRN 611233-15-9
     CMF C21 H18 C12 Y
     CCI CCS
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CRN 444121-94-2 CMF C16 H34 Li O4 CCI CCS

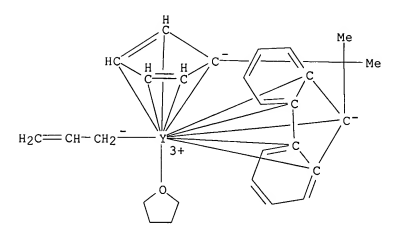
IT 714977-58-9P

RL: CAT (Catalyst use); IMF (Industrial manufacture); PREP (Preparation); USES (Uses)

(metallocenes bridged with Group IIIB elements and based on cyclopentadienyl-fluorenyl ligands for catalysts for polymerization of polar and nonpolar monomers)

RN 714977-58-9 HCAPLUS

CN Yttrium, [n10-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene]-2-propenyl(tetrahydrofuran)- (9CI) (CA INDEX NAME)



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Referenced Author	Year VC	/L) (RPG)	Referenced Work	Referenced
(RAU)	(RPY) (RV		(RWK)	File
Yasuda, H	1995	•	JP 07258319 A	HCAPLUS

L70 ANSWER 5 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

AN 2004:354982 HCAPLUS

DN 140:340585

- TI Ethylene/butadiene copolymers, catalytic system of producing same and production of said polymers
- IN Monteil, Vincent; Spitz, Roger; Boisson, Christophe
- PA Societe De Technologie Michelin, Fr.; Michelin Recherche Et Technique S.A.; Atofina Research
- SO PCT Int. Appl., 31 pp.

CODEN: PIXXD2

DT Patent

LA French

FAN.CNT 1

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PRAI FR 2002-12893 A 20021016 <-- WO 2003-EP11303 W 20031013 <--

OS MARPAT 140:340585

AB The copolymers comprise a molar ratio of units derived from butadiene which is greater than or equal to 8 %, said units comprising trans-1,2-cyclohexane chain formations, and have an number-average mol. weight which

is greater than or equal to 40,000 g/mol. The catalytic system consists of (i) an organometallic complex which is represented by Cp1(Cp2)LnX (I) wherein: Ln represents a lanthanide, X represents a halogen, and Cp1 and Cp2 are each formed by a fluorenyl group, or an organometallic complex composed of similar components as I but Cp1 and Cp2 are also bridged by MR2, wherein M is an element from column IVA and R is an alkyl with between 1 and 20 carbon atoms. The catalytic system also consists of (ii) a co-catalyst belonging to the group comprising a magnesium alkyl, a lithium alkyl, an aluminum alkyl, a Grignard reagent or a mixture of said components. Said catalytic system is such that the molar ratio (co-catalyst/organometallic complex) lies between 1 and 8.

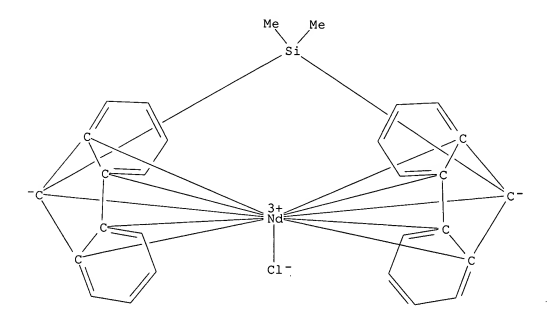
IT 334834-50-3P

RL: CAT (Catalyst use); IMF (Industrial manufacture); PREP (Preparation); USES (Uses)

(manufacture of ethylene-butadiene copolymers in presence of metallocene complexes of lanthanide metals and fluorenyl groups optionally bridged by Group IVA element-based groups)

RN 334834-50-3 HCAPLUS

CN Neodymium, chloro[(dimethylsilylene)bis[(4a,4b,8a,9,9a-η)-9H-fluoren-9ylidene]]- (9CI) (CA INDEX NAME)



RETABLE

Referenced Author (RAU)	(RPY) (RVL) (R	-, ,	Referenced File
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Cui, L Evans, W Llauro, M Michelin Rech Tech	1998 40 72 1994 27 40 2001 34 63 2001	11 MACROMOLECULES	HCAPLUS HCAPLUS HCAPLUS HCAPLUS

Nakamura, H | 2000 | 19 | 5392 | ORGANOMETALLICS | HCAPLUS

L70 ANSWER 6 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

AN 2003:761605 HCAPLUS

DN 139:381567

- TI "Constrained Geometry" Group 3 Metal Complexes of the Fluorenyl-Based Ligands [(3,6-tBu2Flu)SiR2NtBu]: Synthesis, Structural Characterization, and Polymerization Activity
- AU Kirillov, Evgueni; Toupet, Loic; Lehmann, Christian W.; Razavi, Abbas; Carpentier, Jean-Francois
- CS Organometalliques et Catalyse, Groupe Matiere Condensee et Materiaux Cristallochimie, Universite de Rennes 1, UMR 6509 CNRS, UMR 6626 CNRS, Rennes, 35042, Fr.
- SO Organometallics (2003), 22(22), 4467-4479 CODEN: ORGND7; ISSN: 0276-7333
- PB American Chemical Society
- DT Journal
- LA English
- OS CASREACT 139:381567
- Alkane elimination between Y(CH2SiMe3)3(THF)2 and the diprotio ligands AB [(3,6-tBu2C13H7)SiR2NHtBu] (R = Me, la; R = Ph, lb) gave $[\eta 3:\eta 1-((3,6-tBu2C13H6)SiR2NtBu)Y(CH2SiMe3)(THF)2]$ (R = Me, 2a; R = Ph, 2b). 2A is thermally stable in toluene solution and shows a dynamic behavior connected to THF dissociation, while 2b is thermally unstable. Reaction of 2a with H2 or PhSiH3 led to the putative hydrido complex "[(3,6-tBu2Flu)(SiMe2NtBu)YH(THF)]n" (3). Deprotonation of 1a with 1 and 2 equivalent of nBuLi gave [(3,6-tBu2C13H6)SiMe2NHtBu]Li (5) and [(3,6-tBu2C13H6)SiMe2NtBu]Li2 (4), resp., both of which were characterized crystallog. Salt elimination reactions between LnCl3(THF)n precursors (Ln = Y, La, Nd) and 1 equivalent of 4 gave mixts. of complexes, from which ionic complexes that contain two chelated ligands per lanthanide center, $[\eta 3:\eta 1-(3,6-tBu2C13H6)SiMe2NtBu]2Ln]-[Li(solvent)n]+(Ln = Y,$ solvent = THF, n = 4, 6; Ln = La, solvent = THF, n = 4, 7; Ln = La, solvent = Et2O, n = 2, 8; Ln = Nd, solvent = THF, n = 4, 9), were isolated. The neutral dimeric chloro complex $[\eta 5:\eta 1-((3,6$ tBu2C13H6) SiMe2NtBu) Nd(μ -Cl) (THF)]2 (10) was also crystallized from the crude metathesis product. The solid-state structures of 2a, 8, 9, and 10 show versatile coordination modes of the fluorenyl ligands, either $\eta 3$ or $\eta 5$ sym., involving carbon atoms of the central Cp ring (8 and 10), or unusual $\eta 3$ dissym., involving carbon atoms of the central Cp and one adjacent Ph rings (2a and 9). Some of the complexes obtained were explored as catalysts for ethylene and MMA polymerization
- IT 624739-68-0P 624739-71-5P 625094-77-1P
- RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (crystal structure; preparation, crystal structure, and polymerization catalytic
 - activity of constrained geometry lanthanide complexes of aminosilyl fluorenyl-based ligands)
- RN 624739-68-0 HCAPLUS
- CN Lithium, $[\mu-[1-[(8a,9,9a-\eta)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl-\kappaC9]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κN]][[1-[(8a,9,9a-\eta)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]- N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κN]lanthanum]bis[1,1'-oxybis[ethane]]- (9CI) (CA INDEX NAME)$

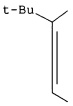
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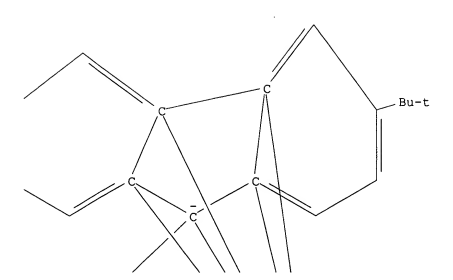
RN 624739-71-5 HCAPLUS

CN Neodymium, bis[1-[(4a,4b,8a,9,9a- η)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κ N]di- μ -chlorobis(tetrahydrofuran)di-, stereoisomer (9CI) (CA INDEX NAME)

PAGE 1-A

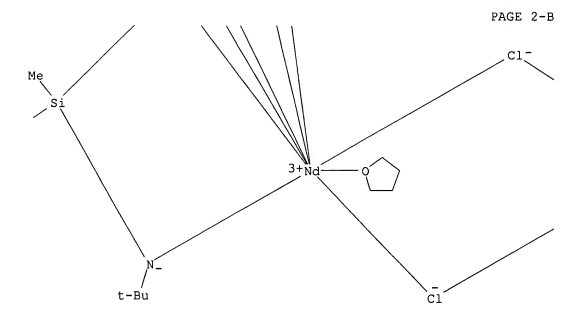


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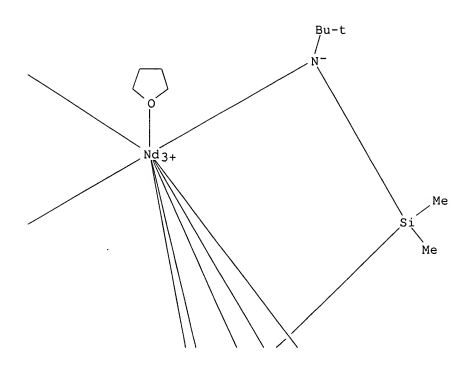


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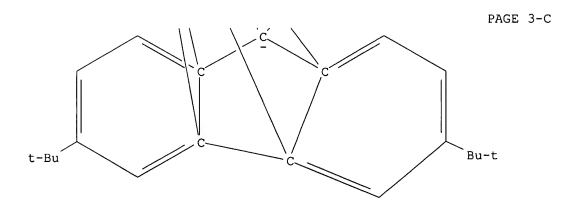
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PAGE 2-C



PAGE 3-A



RN 625094-77-1 HCAPLUS

Lithium(1+), tetrakis(tetrahydrofuran)-, (T-4)-, stereoisomer of $[1-[(1,9,9a-\eta)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)-<math>\kappa$ N][1-[(8a,9- η)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κ N]neodymate(1-), compd. with methylbenzene and tetrahydrofuran (1:1:1) (9CI) (CA INDEX NAME)

CM 1

CRN 109-99-9 CMF C4 H8 O



CM 2

CRN 108-88-3 CMF C7 H8



CM 3

CRN 624739-70-4 CMF C58 H86 N2 Nd O Si2 . C16 H32 Li O4

CM 4

CRN 624739-69-1 CMF C58 H86 N2 Nd O Si2 CCI CCS

PAGE 1-A

PAGE 2-A

CM 5

CRN 48186-27-2 CMF C16 H32 Li O4

CCI CCS

IT 624739-70-4P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (mol. structure; preparation, crystal structure, and polymerization catalytic activity of constrained geometry lanthanide complexes of aminosilyl fluorenyl-based ligands)

RN 624739-70-4 HCAPLUS

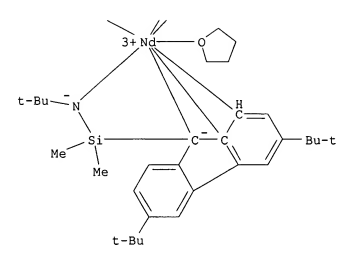
CN Lithium(1+), tetrakis(tetrahydrofuran)-, (T-4)-, stereoisomer of $[1-[(1,9,9a-\eta)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)-<math>\kappa$ N][1-[(8a,9- η)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κ N]neodymate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 624739-69-1 CMF C58 H86 N2 Nd O Si2 CCI CCS

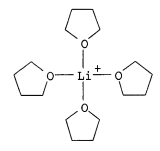
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PAGE 2-A



CM 2

CRN 48186-27-2 CMF C16 H32 Li O4 CCI CCS



IT 624739-61-3P 624739-67-9P

RL: CAT (Catalyst use); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

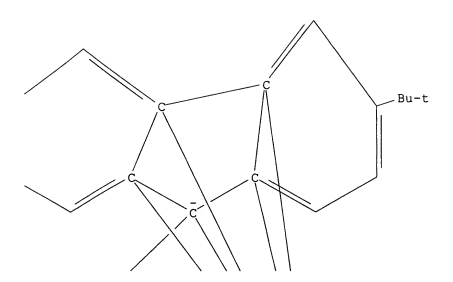
(preparation, crystal structure, and polymerization catalytic activity of constrained geometry lanthanide complexes of aminosilyl fluorenyl-based ligands)

RN 624739-61-3 HCAPLUS

CN Yttrium, bis[1-[(4a,4b,8a,9,9a- η)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κ N]di- μ -hydrobis(tetrahydrofuran)di- (9CI) (CA INDEX NAME)

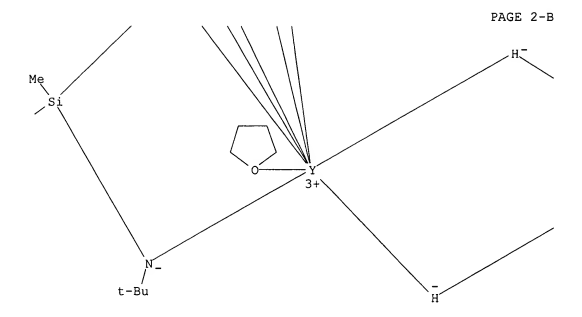
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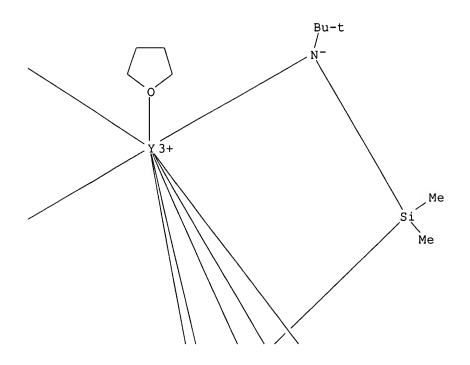


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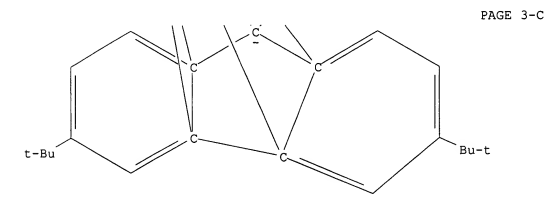
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PAGE 2-C



PAGE 3-A



RN 624739-67-9 HCAPLUS

CN Lithium(1+), tetrakis(tetrahydrofuran)-, (T-4)-, bis[1-[(8a,9,9a- η)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κ N]lanthanate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 624739-66-8

CMF C54 H78 La N2 Si2

CCI CCS

CM 2

CRN 48186-27-2 CMF C16 H32 Li O4 CCI CCS

IT 624739-65-7P

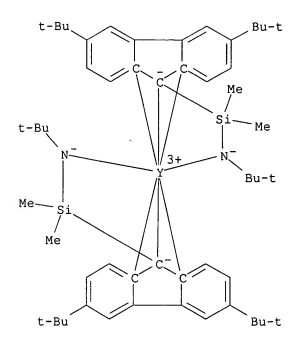
RL: SPN (Synthetic preparation); PREP (Preparation) (preparation, crystal structure, and polymerization catalytic activity of constrained geometry lanthanide complexes of aminosilyl fluorenyl-based ligands)

RN 624739-65-7 HCAPLUS

CN Lithium(1+), tetrakis(tetrahydrofuran)-, (T-4)-, bis[1-[(8a,9,9a- η)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κ N]yttrate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 624739-64-6 CMF C54 H78 N2 Si2 Y CCI CCS



CM 2

CRN 48186-27-2 CMF C16 H32 Li O4 CCI CCS

RETABLE

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Bogaert, S	2001	20	199	Organometallics	HCAPLUS
Boisson, C	12000		75	Progress and Develop	
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Evans, W	1994		1281	Organometallics	HCAPLUS
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Giardello, M	1995	117	3276	J Am Chem Soc	HCAPLUS
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Hultzsch, K	1999	38	227	Angew Chem, Int Ed	HCAPLUS
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Nishiura, M			11184		HCAPLUS
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Okuda, J	•		351		HCAPLUS
Piers, W	1990	•	74	Synlett	HCAPLUS
Qian, C	1999		3283	J Chem Soc, Dalton T	HCAPLUS
Qian, C	2001	626	171	J Organomet Chem	HCAPLUS
Qian, C	12002	645	82	J Organomet Chem	HCAPLUS
Qian, C	12000	119	4134	Organometallics	HCAPLUS
Razavi, A			267		HCAPLUS
Razavi, A	2001		267	Organometallic Catal	
Roesky, P		•	14756	-	HCAPLUS
-			3091	-	HCAPLUS
Ryu, J				-	
Schmid, M			13	_	HCAPLUS
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Shannon, R			751		HCAPLUS
Shapiro, P	11994	116	4623	J Am Chem Soc	HCAPLUS

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Yoder, J
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L70 ANSWER 7 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

AN 2003:664027 HCAPLUS

DN 139:307861

TI [(Cp-CMe2-Flu)2Ln]-[Li(ether)n]+ (Ln = Y, La): Complexes with Unusual Coordination Modes of the Fluorenyl Ligand and the First Examples of Bis-Ansa Lanthanidocenes

AU Kirillov, Evgueni; Toupet, Loic; Lehmann, Christian W.; Razavi, Abbas; Kahlal, Samia; Saillard, Jean-Yves; Carpentier, Jean-Francois

- CS Organometalliques et Catalyse, Institut de Chimie de Rennes, UMR 6509 CNRS-Universite de Rennes 1, Rennes, 35042, Fr.
- SO Organometallics (2003), 22(20), 4038-4046 CODEN: ORGND7; ISSN: 0276-7333
- PB American Chemical Society
- DT Journal
- LA English
- OS CASREACT 139:307861
- Salt metathesis reactions between LnCl3(THF)n (Ln = Y, La) and 1 equivalent of AΒ the dilithium salt of the isopropylidene-bridged ligand [Flu-CMe2-Cp]Li2 (Flu = fluorenyl), in di-Et ether solution, led to the isolation of new ionic metallocene complexes, [(Cp-CMe2-Flu)2Ln]-[Li(ether)n]+ (ether = Et20, THF; Ln = Y, n = 4, 2; Ln = La, n = 2, 3), which contain two chelating ligand units per metal center. The ionic complex 2 presumably originates from ligand redistribution in the primary formed heteroleptic ate complex [(Cp-CMe2-Flu)YCl2]-[Li(ether)4]+ (1) upon crystallization Complex 2 was selectively prepared on using 2 equivalent of [Cp-CMe2-Flu]Li2 vs YCl3(THF)3.5. The solid-state structures of 2 and 3 were established by x-ray diffraction studies. Three polymorphic varieties of 2 were identified and all shown to correspond to a fully dissociated ion pair with the formula $[(\eta 3:\eta 5-Flu-CMe2-Cp)(\eta 1:\eta 5-Flu-CMe2-Cp)Y]-$ [Li(Et2O)(THF)3]+ (2). The fluorenyl ligands in 2 show an unprecedented $\eta 1$ bonding mode and a rare $\eta 3$ bonding mode involving, resp., a carbon atom of a Ph ring and the bridgehead carbon atom of the central ring and the two adjacent carbon atoms of one six-membered ring. DFT computations carried out on the anionic fragment of 2 corroborated the nature of these bonding modes. Only the last exocyclic $\eta 3$ -bonding mode is observed for the fluorenyl moieties in complex 3, which features an associated ion-pair structure with the formula [(η 3: η 5-Flu-CMe2-Cp)2La]-[Li(OEt2)2]+. For comparison purposes, the isopropylidene-bridged bis(indenyl) complex [(Ind-CMe2-Ind)2Y]-[Li(THF)4]+ (4) was synthesized by a salt metathesis procedure and characterized by x-ray diffraction. In contrast to fluorenyl-containing complexes 2 and 3, only the cyclopentadienyl rings of the indenyl moieties coordinate to yttrium in 4. Ionic complexes 2-4 constitute the first structurally characterized examples of bis-ansa lanthanidocenes.
- IT 611233-17-1

RL: PRP (Properties)

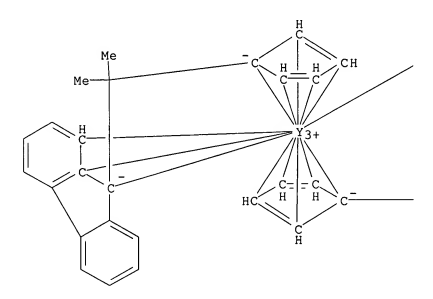
(DFT calcn.; preparation, crystal structure, and DFT studies of isopropylidene bridged fluorenyl cyclopentadienyl lanthanide complexes with unusual coordination modes of fluorenyl ligand)

RN 611233-17-1 HCAPLUS

CN

Yttrate(1-), rel-[η 6-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-(1R)-1H-fluorene-1,9-diyl][η 8-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-(1R)-9H-fluoren-9-ylidene]- (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 1-B

IT 611233-18-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (crystal structure, monoclinic and orthorhombic polymorphs; preparation,

crystal structure, and DFT studies of isopropylidene bridged fluorenyl cyclopentadienyl lanthanide complexes with unusual coordination modes of fluorenyl ligand)

RN 611233-18-2 HCAPLUS CN Lithium(1+), [1,1'-o:

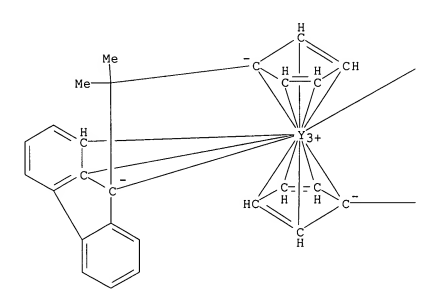
Lithium(1+), [1,1'-oxybis[ethane]]tris(tetrahydrofuran)-, (T-4)-, stereoisomer of [η6-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-1H-fluorene-1,9-diyl][η8-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene]yttrate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 611233-17-1 CMF C42 H36 Y

CCI CCS

PAGE 1-A



PAGE 1-B

CM 2

CRN 444121-94-2 CMF C16 H34 Li O4 CCI CCS

IT 612060-80-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(crystal structure, triclinic polymorph; preparation, crystal structure, and DFT studies of isopropylidene bridged fluorenyl cyclopentadienyl lanthanide complexes with unusual coordination modes of fluorenyl ligand)

RN 612060-80-7 HCAPLUS

CN Lithium(1+), [1,1'-oxybis[ethane]]tris(tetrahydrofuran)-, (T-4)-, stereoisomer of [η 6-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-1H-fluorene-9,1-diyl][η 8-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene]yttrate(1-), compd. with 1,1'-oxybis[ethane] (2:1) (9CI) (CA INDEX NAME)

CM 1

CRN 60-29-7 CMF C4 H10 O

H3C-CH2-O-CH2-CH3

CM 2

CRN 611233-18-2

CMF C42 H36 Y . C16 H34 Li O4

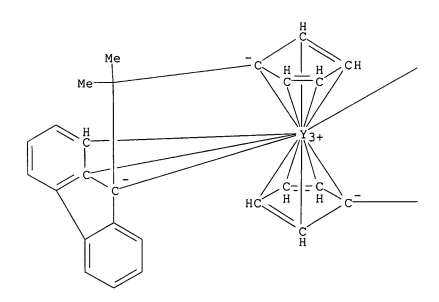
CM 3

CRN 611233-17-1

CMF C42 H36 Y

CCI CCS

PAGE 1-A



PAGE 1-B

CM 4

CRN 444121-94-2 CMF C16 H34 Li O4 CCI CCS

IT 611233-16-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

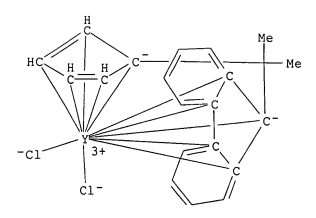
(preparation, crystal structure, and DFT studies of isopropylidene bridged fluorenyl cyclopentadienyl lanthanide complexes with unusual coordination modes of fluorenyl ligand)

RN 611233-16-0 HCAPLUS

CN Lithium(1+), [1,1'-oxybis[ethane]]tris(tetrahydrofuran)-, (T-4)-, dichloro[\eta10-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene]yttrate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 611233-15-9 CMF C21 H18 C12 Y CCI CCS



CM 2

CRN 444121-94-2 CMF C16 H34 Li O4

CCI CCS

RETABLE

Referenced Author (RAU)	Year VOL (RPY) (RVL) [(RPG)	Referenced Work (RWK)	Referenced File
Alt, H Anwander, R Anwander, R Anwander, R Arndt, S Baerends, E Baerends, E Becke, A Bickelhaupt, F Bochmann, M Bochmann, M Bochmann, M Boerringter, P Brintzinger, H Coates, G Dash, A Den Haan, K Drago, D Drago, D Edelmann, F	1996 100 1996 2 1996 179 1999 2 1973 12 1988 A38 2000 15 1996 11988 33 1995 34 12000 12002 21 1986 5 12000 19 12000 19 1996 179	1205 866 1 1953 41 169 3098 1 255 4718 87 1143 1223 3238 1726 1802 1802 247	Chem Rev Applied Homogeneous Top Curr Chem Top Organomet Chem Top Organomet Chem Chem Rev Chem Phys Int J Quantum Chem Phys Rev Rev Comput Chem J Chem Soc, Dalton Organometallics Int J Quantum Chem Angew Chem, Int Ed F Chem Rev Organometallics Organometallics Organometallics Organometallics Organometallics Organometallics Organometallics Organometallics Organometallics Top Curr Chem	HCAPLUS HCAPLUS HCAPLUS HCAPLUS HCAPLUS HCAPLUS HCAPLUS HCAPLUS
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L70 ANSWER 8 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN
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AN 2002:570105 HCAPLUS

DN 138:271764

TI Synthesis and characterization of organolanthanide chlorides Me2SiFlu2LnCl

AU Zhang, Wu; Cai, Yue-peng; Li, Hong-xi; Ma, Huai-zhu

CS College of Chemistry and Material Science, Anhui Normal University, Wuhu, 241000, Peop. Rep. China

SO Hecheng Huaxue (2002), 10(3), 268-270 CODEN: HEHUE2; ISSN: 1005-1511

PB Hecheng Huaxue Bianjibu

DT Journal

LA Chinese

OS CASREACT 138:271764

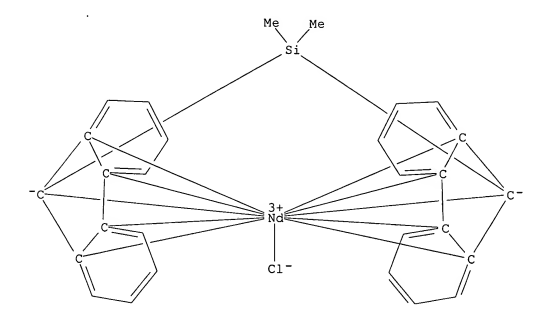
AB Five new silyl-bridged organolanthanide complexes Me2SiFlu2LnCl (Flu = fluorenyl, Ln = Yb, Sm, La, Pr, Nd) were synthesized by the reaction of LnCl3 with Me2SiFlu2Li2 in THF. These complexes were characterized by elemental analyses, IR, MS and 1H NMR spectra.

IT 334834-50-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (synthesis and characterization of silyl-bridged fluorenyl lanthanide chlorides)

RN 334834-50-3 HCAPLUS

CN Neodymium, chloro[(dimethylsilylene)bis[(4a, 4b, 8a, 9, 9a-η)-9H-fluoren-9ylidene]]- (9CI) (CA INDEX NAME)



- L70 ANSWER 9 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN
- AN 2002:457779 HCAPLUS
- DN 137:169594
- TI Amine Elimination Reactions between Homoleptic Silylamide Lanthanide Complexes and an Isopropylidene-Bridged Cyclopentadiene-Fluorene System
- AU Dash, Aswini K.; Razavi, Abbas; Mortreux, Andre; Lehmann, Christian W.; Carpentier, Jean-Francois
- CS Laboratoire Organometalliques et Catalyse, UMR 6509 CNRS-Universite de Rennes 1, Rennes, 35042, Fr.
- SO Organometallics (2002), 21(15), 3238-3249 CODEN: ORGND7; ISSN: 0276-7333
- PB American Chemical Society
- DT Journal
- LA English
- OS CASREACT 137:169594
- Amine elimination to introduce the isopropylidene-bridged unsym. ligand C5H5-CMe2-C13H9 (CpH-CMe2-FluH) onto group III-metal centers (Y, La, Nd) to give the neutral, ate-complex-free ansa-lanthanidocenes is discussed. The reactions of homoleptic Ln[N(SiMe3)2]3 (Ln = Y (1), La (2), Nd (3)) with CpH-CMe2-FluH (4) in THF under mild conditions lead to the formation of ansa-complexes (η5,η5-Cp-CMe2-Flu)Ln(η5-Cp-CMe2-FluH) (Ln = Y (8), La (12), Nd (13)) in 70-84% isolated yields (based on 4). These reactions proceed via the rapidly formed bis(amido)lanthanide intermediates (η5-Cp-CMe2-FluH)Ln[N(SiMe3)2]2 (Ln = Y (5), La (9)), which undergo readily disproportionation/ligand redistribution reactions at 5-23° to give either a mono(amido)lanthanide complex (η5-Cp-CMe2-FluH)2Ln[N(SiMe3)2] (Ln = Y (6)) or another species

assumed to be the binuclear complex (η5-Cp-CMe2-FluH)2Ln[μ-N(SiMe3)2]2Ln[N(SiMe3)2]2 (Ln = La (10)), resp. Complexes 6 and 10 undergo an intramol. amine elimination reaction under THF reflux to yield the corresponding ansa-complexes 8 and 12, resp. The reversibility of the process was investigated in the yttrium case: complex 8 converts back to 6 in the presence of (SiMe3)2NH in toluene at 90° with 50% conversion after 12 h. The effect of a noncoordinating apolar solvent on the reaction outcome of tris(amido) complexes 1-3 with 4 was also studied using toluene, in which the low solubility presumably shifts the disproportionation equilibrium and leads to the isolation of another class of compds. Ln(η 5-Cp-CMe2-FluH)3 (Ln = Y (7), La (11)) in reasonable yields. Compds. 5-12 were characterized in solution by 1D and 2D NMR techniques (1H, 13C, 1H-1H COSY, and 1H-13C HETCOR), and the solid state structures of 6 and of the mono(THF) adducts of ansa-lanthanidocenes 12and 13 were established by x-ray diffraction studies. The latter ansa-complexes feature very narrow Cp(centroid)-Ln-Flu(centroid) bite angles (Ln = La, $103.67(1)^{\circ}$; Ln = Nd, $105.08(1)^{\circ}$).

IT 447452-98-4P 447453-02-3P 447453-03-4P 447453-05-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(amine elimination reactions between homoleptic silylamide lanthanide complexes and isopropylidene-bridged cyclopentadiene-fluorene system) 447452-98-4 HCAPLUS

Yttrium, $[(1,2,3,4,5-\eta)-1-[1-(9H-fluoren-9-yl)-1-methylethyl]-2,4-$ cyclopentadien-1-yl]bis[1,1,1-trimethyl-N-(trimethylsilyl)silanaminato]-(9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ &$$

RN

CN

RN 447453-02-3 HCAPLUS

CN Yttrium, [η10-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9Hfluoren-9-ylidene][(1,2,3,4,5-η)-1-[1-(9H-fluoren-9-yl)-1-methylethyl]2,4-cyclopentadien-1-yl]- (9CI) (CA INDEX NAME)

RN 447453-03-4 HCAPLUS

CN Lanthanum, $[(1,2,3,4,5-\eta)-1-[1-(9H-fluoren-9-yl)-1-methylethyl]-2,4-cyclopentadien-1-yl]bis[1,1,1-trimethyl-N-(trimethylsilyl)silanaminato]-(9CI) (CA INDEX NAME)$

$$\begin{array}{c|c} & H & C & C & Me \\ \hline H & C & C & Me \\ \hline La^{3+} & N-SiMe_3 \\ \hline SiMe_3 & SiMe_3 \\ \end{array}$$

RN 447453-05-6 HCAPLUS

CN Lanthanum, bis[$(1,2,3,4,5-\eta)-1-[1-(9H-fluoren-9-yl)-1-methylethyl]-2,4-cyclopentadien-1-yl]$ bis[μ -[1,1,1-trimethyl-N-(trimethylsilyl)silanaminato]]bis[1,1,1-trimethyl-N-(trimethylsilyl)silanaminato]di-(9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

IT 447452-99-5DP, agostic bond

RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (crystal structure; amine elimination reactions between homoleptic

silylamide lanthanide complexes and isopropylidene-bridged cyclopentadiene-fluorene system)

RN 447452-99-5 HCAPLUS

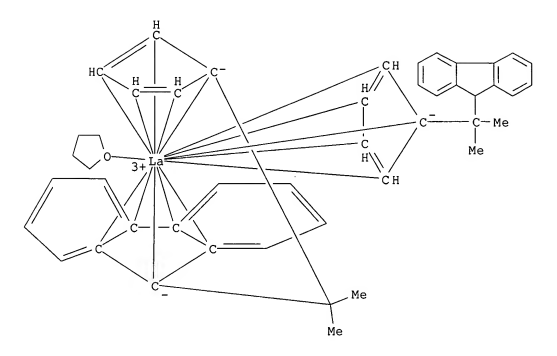
CN Yttrium, bis $[(1,2,3,4,5-\eta)-1-[1-(9H-fluoren-9-yl)-1-methylethyl]-2,4-cyclopentadien-1-yl][1,1,1-trimethyl-N-(trimethylsilyl)silanaminato]-(9CI) (CA INDEX NAME)$

IT 447453-17-0P 447453-21-6P 447453-25-0P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and crystal structure of)

RN 447453-17-0 HCAPLUS

CN Lanthanum, $[\eta 10-2, 4-\text{cyclopentadien}-1-\text{ylidene}(1-\text{methylethylidene})-9H-fluoren-9-ylidene][(1,2,3,4,5-<math>\eta$)-1-[1-(9H-fluoren-9-yl)-1-methylethyl]-2,4-cyclopentadien-1-yl](tetrahydrofuran)- (9CI) (CA INDEX NAME)



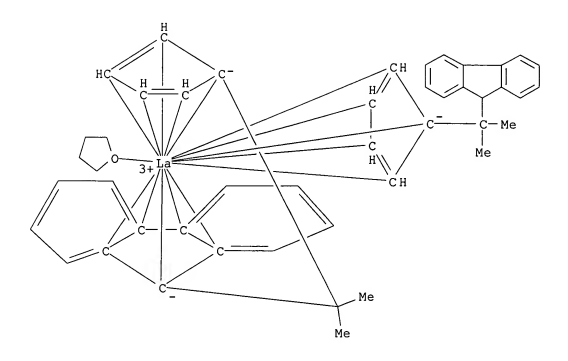
RN447453-21-6 HCAPLUS

Lanthanum, [η 10-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-CN fluoren-9-ylidene] $[(1,2,3,4,5-\eta)-1-[1-(9H-fluoren-9-yl)-1-methylethyl]-1$ 2,4-cyclopentadien-1-yl](tetrahydrofuran)-, compd. with tetrahydrofuran (2:3) (9CI) (CA INDEX NAME)

CM 1

CRN 447453-17-0 CMF C46 H45 La O

CCI CCS



CM 2

CRN 109-99-9 CMF C4 H8 O



CN

RN 447453-25-0 HCAPLUS

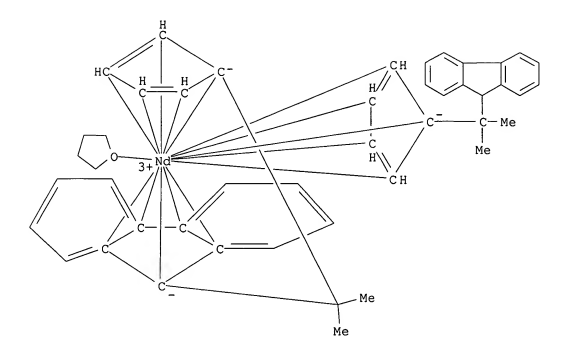
Neodymium, $[\eta 10-2, 4-\text{cyclopentadien}-1-\text{ylidene}(1-\text{methylethylidene})-9H-fluoren-9-\text{ylidene}][(1,2,3,4,5-\eta)-1-[1-(9H-fluoren-9-\text{yl})-1-\text{methylethyl}]-2,4-\text{cyclopentadien}-1-\text{yl}](\text{tetrahydrofuran})-, \text{compd. with tetrahydrofuran}(1:2) (9CI) (CA INDEX NAME)$

CM 1

CRN 447453-12-5

CMF C46 H45 Nd O

CCI CCS



CM 2

CRN 109-99-9 CMF C4 H8 O



447453-00-1P 447453-08-9P 447453-10-3P IT

447453-12-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

447453-00-1 HCAPLUS RN

Yttrium, tris $[(1,2,3,4,5-\eta)-1-[1-(9H-fluoren-9-yl)-1-methylethyl]-2,4-$ CN cyclopentadien-1-yl]- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

RN 447453-08-9 HCAPLUS CN Lanthanum, tris[$(1,2,3,4,5-\eta)-1-[1-(9H-fluoren-9-yl)-1-methylethyl]-2,4-cyclopentadien-1-yl]- (9CI) (CA INDEX NAME)$

PAGE 1-A

$$\begin{array}{c} H \\ R \\ H \\ C \\$$

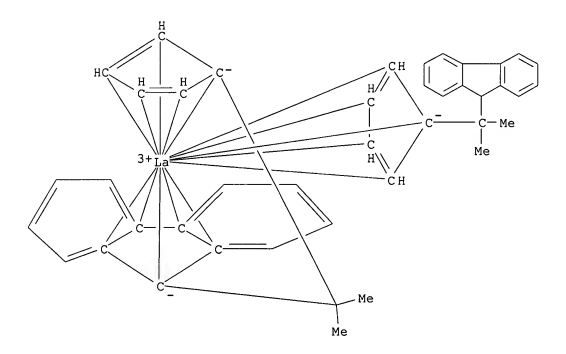
PAGE 2-A

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PAGE 3-A

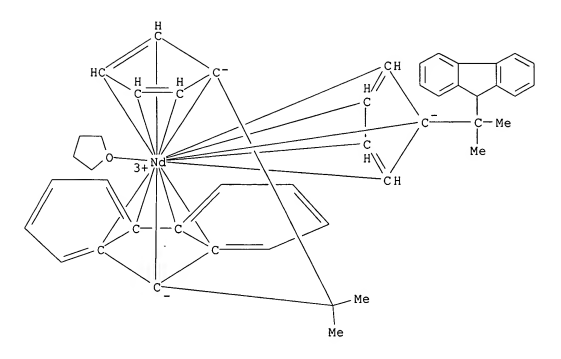
RN 447453-10-3 HCAPLUS

CN Lanthanum, $[\eta 10-2, 4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene][(1,2,3,4,5-<math>\eta$)-1-[1-(9H-fluoren-9-yl)-1-methylethyl]-2,4-cyclopentadien-1-yl]- (9CI) (CA INDEX NAME)



RN 447453-12-5 HCAPLUS

CN Neodymium, $[\eta 10-2, 4-\text{cyclopentadien}-1-\text{ylidene}(1-\text{methylethylidene})-9\text{H-fluoren}-9-\text{ylidene}][(1,2,3,4,5-\eta)-1-[1-(9\text{H-fluoren}-9-\text{yl})-1-\text{methylethyl}]-2,4-\text{cyclopentadien}-1-\text{yl}](tetrahydrofuran)- (9CI) (CA INDEX NAME)$



RETABLE					
Referenced Author	Year	VOL	PG	Referenced Work	Referenced
(RAU)	(RPY)	(RVL)	(RPG)	(RWK)	File
	+=====	+=====	+=====		+=======
Akhnoukh, T		•	47	. 3	HCAPLUS
Alt, H	•		1205		HCAPLUS
Alt, H	1998	27	323	•	HCAPLUS
Anwander, R		12	1866		HCAPLUS
Anwander, R		179	1	Top Curr Chem	HCAPLUS
Anwander, R	•		133	Top Curr Chem	HCAPLUS
Anwander, R	1999	2	1	Top Organomet Chem	HCAPLUS
Avent, A	11989	111	3423	J Am Chem Soc	HCAPLUS
Ballard, D	1978	1	1994	J Chem Soc, Chem Com	HCAPLUS
Barbier-Baudry, D	11998	1	1721	Eur J Inorg Chem	l
Barbier-Baudry, D	12000	609	21	•	HCAPLUS
Bochmann, M	1996	1	255	J Chem Soc, Dalton T	HCAPLUS
Bogaert, S	12000	201	1813	Macromol Chem Phys	HCAPLUS
Bogaert, S	2001	20	199	, 3	HCAPLUS
Boncella, J	1985	4	1205		HCAPLUS
Booj, M	1989	364	79	J Organomet Chem	l
Bordwell, F	11983	105	6188	J Am Chem Soc	HCAPLUS
Bradley, D	1972		349	J Chem Soc, Chem Com	HCAPLUS
Bradley, D	1973	1	1021	J Chem Soc, Dalton T	HCAPLUS
Brintzinger, H	•	34	1143	Angew Chem, Int Ed E	HCAPLUS
Britovsek, G	1999	138	428		HCAPLUS
Burger, B			1566	J Am Chem Soc	HCAPLUS
Chen, E		100	1391	•	HCAPLUS
Coughlin, E	•	1114	17606	• • • • • • • • • • • •	HCAPLUS
Coughlin, E	•	133	11226	Polym Prepr, Am Chem	
Den Haan, K		5	1726	Organometallics	HCAPLUS
Drago, D	•	19	1802	Organometallics	HCAPLUS
Edelmann, F	•		1247	Top Curr Chem	HCAPLUS
Eppinger, J	•	122	13080	J Am Chem Soc	HCAPLUS
Evans, W	1983	1104	12008	J Am Chem Soc	

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Evans, W	1990	112	2314	J Am Chem Soc	HCAPLUS
Evans, W	1985	4	554	Organometallics	HCAPLUS
Evans, W	1993	112			HCAPLUS
	1994			· -	HCAPLUS
•		-	6255		HCAPLUS
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	•		174		HCAPLUS
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			4134	=	HCAPLUS
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	1993		117		HCAPLUS
· ·			267	-	HCAPLUS
-			1383	J Organomet Chem	
•			1602		HCAPLUS
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The state of the s		-	751	Acta Crystallogr, Se	HCAPLUS
·		•	1925	Acta Crystallogr, Se	
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= '		-	4623	J Am Chem Soc	HCAPLUS
-	•	9	1867	Organometallics	HCAPLUS
	1997		1	SHELXL-97, Program f	
	1997	1		SHELXS-97, Program f	
	•		12568	Organometallics	HCAPLUS
- '	1984	23	2271	Inorg Chem	HCAPLUS
van der Heijden, H	1989	8	1255	Organometallics	HCAPLUS
			459	ACS Symp Ser	HCAPLUS
Watson, P	1985		51		HCAPLUS
Westernhausen, M	•		837	Z Anorg Allg Chem	
Yan, P	1990	391	313		HCAPLUS

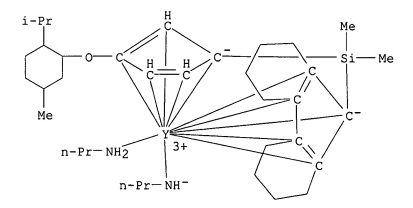
- L70 ANSWER 10 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN 2001:912364 HCAPLUS AN DN 136:183902 TΤ "Widening the Roof": Synthesis and Characterization of New Chiral C1-Symmetric Octahydrofluorenyl Organolanthanide Catalysts and Their Implementation in the Stereoselective Cyclizations of Aminoalkenes and Phosphinoalkenes ΑU Douglass, Michael R.; Ogasawara, Masamichi; Hong, Sukwon; Metz, Matthew V.; Marks, Tobin J. CS Department of Chemistry, Northwestern University, Evanston, IL, 60208-3113, USA Organometallics (2002), 21(2), 283-292 SO CODEN: ORGND7; ISSN: 0276-7333 PB American Chemical Society DT Journal LA English OS CASREACT 136:183902 AB New chiral C1-sym. organolanthanide ansa-metallocene catalysts Me2Si(OHF)(CpR*)LnN(TMS)2 (OHF = η 5-octahydrofluorenyl; Cp = η 5-C5H3; R* = (-)-menthyl; Ln = Sm, Y, Lu; TMS = SiMe3) were synthesized, characterized, and implemented in the enantioselective and diastereoselective cyclizations of aminoalkenes and phosphinoalkenes. Me2Si(OHF)(CpR*)LnCl2-Li(DME)2+ catalyst precursors can be prepared in up to .apprx.90% diastereomeric purity and then converted into the corresponding amido catalysts, which can be isolated in .apprx.100% diastereomeric purity after recrystn. The catalyst (S)-Me2Si(OHF)(CpR*)YN(TMS)2 was crystallog. characterized. The activity of these catalysts for the hydroamination/cyclization of aminoalkenes and for the hydrophosphination/cyclization of phosphinoalkenes is described. Enantioselectivites ≤67% were obtained in hydroamination, and diastereoselectivities of ≤96% were obtained in hydrophosphination. 400608-55-1P IT RL: CAT (Catalyst use); CPS (Chemical process); PEP (Physical, engineering or chemical process); PRP (Properties); SPN (Synthetic preparation); PREP (Preparation); PROC (Process); USES (Uses) (crystal structure, epimerization mechanism; preparation and catalyst for stereoselective hydroamination or hydrophosphination and cyclization of
- RN 400608-55-1 HCAPLUS
- CN Yttrium, [η10-[(1S)-3-[((1R,2S,5R)-5-methyl-2-(1methylethyl)cyclohexyl]oxy]-2,4-cyclopentadien-1ylidene](dimethylsilylene)(1,2,3,4,5,6,7,8-octahydro-9H-fluoren-9ylidene)][1,1,1-trimethyl-N-(trimethylsilyl)silanaminato]- (9CI) (CA
 INDEX NAME)

aminoalkenes and phosphinoalkenes, resp.)

IT 400608-60-8P

RN 400608-60-8 HCAPLUS

CN Yttrium, [n10-[(1S)-3-[[(1R,2S,5R)-5-methyl-2-(1-methylethyl)cyclohexyl]oxy]-2,4-cyclopentadien-1-ylidene](dimethylsilylene)(1,2,3,4,5,6,7,8-octahydro-9H-fluoren-9-ylidene)](1-propanaminato)(1-propanamine)- (9CI) (CA INDEX NAME)



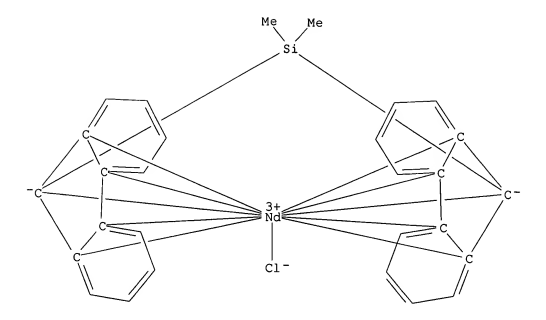
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Baechler, R	1970 92	13090	J Am Chem Soc	HCAPLUS
Bijpost, E	1995 95	121	J Mol Catal	HCAPLUS
Blaser, H	1996 2	1992	Applied Homogeneous	HCAPLUS
Brookhart, M	1988 36	1	Prog Inorg Chem	HCAPLUS
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Coughlin, E	1992	114	7607	J Am Chem Soc	
	1984	120		Phosphorus Sulfur Re	HCAPLUS
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Fu, P	1995	117	7157	J Am Chem Soc	HCAPLUS
Gagne, M	1989	111	4108	J Am Chem Soc	HCAPLUS
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	2001			Abstracts of Papers,	
	1996		2056	J Org Chem	HCAPLUS
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Hultzch, K	1997	18	809	Macromol Rapid Commu	
Hultzch, K	1997	16	4845	Organometallics	
Jacobsen, E	1999	I-II		Comprehensive Asymme	!
Jany, G	1997	126		Organometallics	
	1985		-	_	HCAPLUS
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- '	1999				HCAPLUS
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Molander, G	1998	18	237	Chemtracts:Org Chem	
Molander, G	1992	57	6347	J Org Chem	HCAPLUS
Molander, G	1998	63	8983	J Org Chem	HCAPLUS
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	1994	1		Asymmetric Catalysis	
.		119			HCAPLUS
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The second secon	12000	1 1 0	2503		HCAPLUS
	-	110		Catalytic Asymmetric	
T	•		14134		HCAPLUS
	-		295	The Chemistry of Org	
			400	Angew Chem, Int Ed E	
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Ripperger, H	1965	21	407	Tetrahedron	HCAPLUS

Roesky, P	1997	116	4486	Organometallics HCAPLUS
Ryu, J	2001	13	3091	Abstracts of Papers, HCAPLUS
Sakakura, T	1991	1	40	J Chem Soc, Chem Com HCAPLUS
Schaverien, C	1994	136	283	Adv Organomet Chem HCAPLUS
Schellenberg, J	12000	138	12428	J Polym Sci, Polym C HCAPLUS
Schumann, H	1995	195	865	Chem Rev HCAPLUS
Schumann, H	1998	1559	181	J Organomet Chem HCAPLUS
Shannon, R	1976	A32	751	Acta Crystallogr HCAPLUS
Sheldrick, G	1997	1		SHELXL-97
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Stern, D	1990	1112	9558	J Am Chem Soc HCAPLUS
Watson, P	1982	1104	337	J Am Chem Soc HCAPLUS
Yasuda, H	1995	1196	2417	Macromol Chem Phys HCAPLUS
Yoder, J	1998	17	14946	Organometallics HCAPLUS

- L70 ANSWER 11 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN
- AN 2001:566437 HCAPLUS
- DN 135:273431
- TI Investigation of ethylene/butadiene copolymers microstructure by 1H and 13C NMR
- AU Llauro, Marie France; Monnet, Christiane; Barbotin, Fanny; Monteil, Vincent; Spitz, Roger; Boisson, Christophe
- CS Laboratoire de Chimie et Procedes de Polymerisation, CNRS-CPE Lyon, Villeurbanne, 69616, Fr.
- SO Macromolecules (2001), 34(18), 6304-6311 CODEN: MAMOBX; ISSN: 0024-9297
- PB American Chemical Society
- DT Journal
- LA English
- AB Ethylene and butadiene are copolymd. with neodymocene catalysts. In this paper, a complete 1H and 13C NMR anal. of the copolymers is reported for the first time. The results of this study show that the microstructure depends on the cyclopentadienyl ligands of the catalyst. The presence of the trans-1,2-cyclohexane structure, formed by intramol. cyclization, is detected and fully investigated by 2D NMR 1H/13C direct and long-range correlation.
- IT 334834-50-3
 - RL: CAT (Catalyst use); USES (Uses)
 - (in neodymocene catalysts for polymerization of ethylene with butadiene)
- RN 334834-50-3 HCAPLUS
- CN Neodymium, chloro[(dimethylsilylene)bis[(4a,4b,8a,9,9a- η)-9H-fluoren-9-ylidene]]- (9CI) (CA INDEX NAME)



RETABLE					
Referenced Author	Year	VOL	PG	Referenced Work	Referenced
(RAU)	(RPY)			(RWK)	File
	•	•	•		•
Arnold, M	1991	•	1017	•	HCAPLUS
Barbotin, F	•	33	8521	•	HCAPLUS
Barbotin, F	1999		1	Ph D Thesis, Lyon 1	1
Bruzzone, M	•		2173		HCAPLUS
Cesca, S		•	569	Transition metal cat	•
Desmurs, P	•	•	375	C R Acad Sci Paris I	•
Furukawa, J	-	•	189	Angew Makromol Chem	
Furukawa, J		•	600	Rubber Chem Technol	•
Galimberti, M	1991	192	2591	•	HCAPLUS
Ibbett, R	1993			NMR Spectroscopy of	[
Igai, S	1997		1	JP 9316118	1
Kalinowski, H	1988		112	Carbon-13 NMR Spectr	
Kaminsky, W	•		515	•	HCAPLUS
Kaminsky, W		•	103	Makromol Chem, Macro	HCAPLUS
Kaulbach, R	•	•	101	Angew Makromol Chem	•
Kim, I		•	1590	J Polym Sci A: Polym	
Kudashev, R	•		1398	Dokl Phys Chem	
Moritani, T	•	•	532	•	HCAPLUS
Mulhaupt, R	•	•	2487	J Polym Sci, Part, A	
Naga, N		•	1348	•	HCAPLUS
Natta, G	•	79	161	•	HCAPLUS
Nishiyama, T	1999	l		•	HCAPLUS
Resconi, L	•		4953	•	HCAPLUS
Robert, P			261	Makromol Chem, Macro	
Soga, K	•			, 2	HCAPLUS
Sun, L	•		12113	J Polym Sci, Part B	HCAPLUS
Tiley, T	1981	120	13267	Inorg Chem	1

L70 ANSWER 12 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN AN 2001:279463 HCAPLUS

DN 134:296245

ΤI Catalytic system based on a lanthanide metal complex, process for its preparation and that of an ethylene-conjugated diene copolymer

IN Barbotin, Fanny

PA Societe De Technologie Michelin, Fr.; Michelin Recherche Et Technique S.A.; ATOFINA

SO Eur. Pat. Appl., 26 pp.

CODEN: EPXXDW

DT Patent LA French

FAN.	CNT 1				
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os	MARPAT 134:296245				

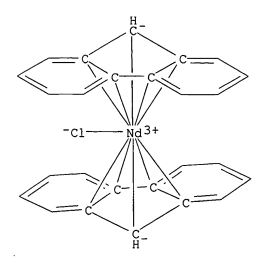
AB Catalysts for manufacture of title copolymers with controlled microstructure of the units formed from the diene contain Cp1(Cp2)LnX [I, Ln = lanthanide metal, X = halo, Cp1, Cp2 = (substituted) cyclopentadienyl or fluorenyl) or a complex similar to I with Cp1 bridged to Cp2 by a Group IVA metal or MR2 (R = C1-20 alkyl, M = Group IVA metal) and cocatalyst selected from alkylmagnesium, alkyllithium, alkylaluminum, or Grignard reagent. A typical I was manufactured by reaction of 20 mmol cyclopentadienyltrimethylsila ne 3 h with 20 mmol BuLi in heptane, and complexation of 8 mmol intermediate 48 h with 4 mmol NdCl3 in THF.

IT 188405-00-7

RL: RCT (Reactant); RACT (Reactant or reagent)
(catalyst precursor; catalysts containing lanthanide metal complexes for
manufacture of ethylene-conjugated diene copolymers)

RN 188405-00-7 HCAPLUS

CN Neodymium, chlorobis[(4a,4b,8a,9,9a-η)-9H-fluoren-9-yl]- (9CI) (CA INDEX NAME)



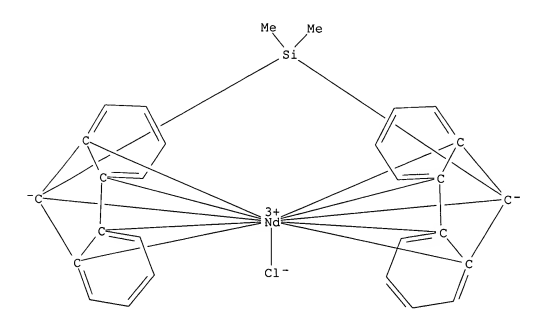
IT 334834-50-3P

RL: CAT (Catalyst use); IMF (Industrial manufacture); PREP (Preparation); USES (Uses)

(catalysts containing lanthanide metal complexes for manufacture of ethylene-conjugated diene copolymers)

RN 334834-50-3 HCAPLUS

CN Neodymium, chloro[(dimethylsilylene)bis[(4a, 4b, 8a, 9, 9a-η)-9H-fluoren-9ylidene]]- (9CI) (CA INDEX NAME)



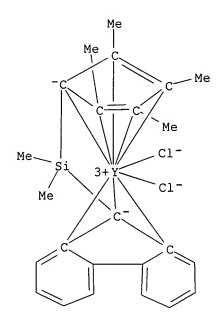
RETABLE

Referenced Author (RAU)	Year VOL (RPY) (RVL) (RPG)	Referenced Work (RWK) =+===========	Referenced File
Cui	1998 40	729	POLYM BULL (BERLIN)	HCAPLUS
Marks, T	1989		US 4801666 A	HCAPLUS
Pettijohn, T	1992		US 5109085 A	HCAPLUS

jan delaval - 4 october 2006

L70 ANSWER 13 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

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AN
     1999:699678 HCAPLUS
DN
     132:64345
ΤI
     The First Fluorenyl ansa-Yttrocene Complexes: Synthesis, Structures, and
     Polymerization of Methyl Methacrylate
     Lee, Min Hyung; Hwang, Jeong-Wook; Kim, Youngjo; Kim, Jindong; Han,
ΑU
     Yonggyu; Do, Youngkyu
CS
     Department of Chemistry and Center for Molecular Science, Korea Advanced
     Institute of Science and Technology, Taejon, 305-701, S. Korea
SO
     Organometallics (1999), 18(24), 5124-5129
     CODEN: ORGND7; ISSN: 0276-7333
PΒ
     American Chemical Society
DT
     Journal
LA
     English
     A novel Cs-sym. yttrocene complex, ansa-Me2Si(η3-Flu)(η5-
AB
     Cp')YCl2Li(OEt2)2 (3; Flu = C13H8, fluorenyl; Cp' = C5Me4), was prepared via
     a salt metathesis reaction from anhydrous YC13 and the dilithium salt of the
     ligand ansa-Me2Si(FluH)(Cp'H). Treatment of 3 with NaN(SiMe3)2 gave the
     corresponding bis(trimethylsilyl)amide derivative ansa-Me2Si(Flu)(η5-
     Cp')YN(SiMe3)2 (4). The x-ray structure of 3 reveals unusual
     \eta 3-fluorenyl coordination to the Y3+ ion. In 4, the Y-Flu bonding
     being partially slipped toward \eta3 from \eta5, \pi-dative bonding
     nature in the Y-N bond and a direct interaction of the Y atom with one Me
     group of the N(SiMe3)2 fragment are present. Both compds. constitute the
     1st examples of structurally characterized fluorenyl ansa-yttrocenes. The
     neutral compound 4 is active for the polymerization of Me methacrylate (MMA) in
     toluene, affording iso-rich poly(MMA)s.
IT
     253305-31-6P
     RL: PEP (Physical, engineering or chemical process); PRP (Properties); SPN
     (Synthetic preparation); PREP (Preparation); PROC (Process)
        (crystal structure; preparation, structure and polymerization catalytic
activity of
        fluorenyl ansa-yttrocene complexes)
RN
     253305-31-6 HCAPLUS
     Lithium(1+), bis[1,1'-oxybis[ethane]]-, dichloro[\eta8-9H-fluoren-9-
CN
     ylidene(dimethylsilylene)(2,3,4,5-tetramethyl-2,4-cyclopentadien-1-
     ylidene)]yttrate(1-) (9CI) (CA INDEX NAME)
     CM
     CRN 253305-30-5
     CMF C24 H26 C12 Si Y
     CCI CCS
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CM 2

CRN 78127-97-6 CMF C8 H20 Li O2 CCI CCS

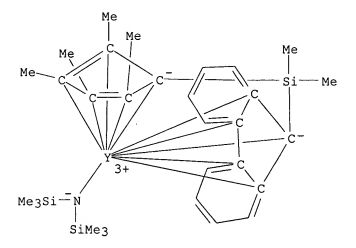
IT 253305-32-7P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (crystal structure; preparation, structure and polymerization catalytic activity of

fluorenyl ansa-yttrocene complexes)

RN 253305-32-7 HCAPLUS

CN Yttrium, [η10-9H-fluoren-9-ylidene(dimethylsilylene) (2,3,4,5-tetramethyl-2,4-cyclopentadien-1-ylidene)][1,1,1-trimethyl-N-(trimethylsilyl)silanaminato]- (9CI) (CA INDEX NAME)



RETABLE Referenced Author (RAU)	Year (RPY) +====	(RVL)	(RPG)	(RWK)	Referenced File +=======
Barnhart, D			8461		HCAPLUS
Brooks, J	11972	94	17339	J Am Chem Soc	HCAPLUS
Chen, Y	11995	1497	1	J Organomet Chem	HCAPLUS
Christopher, J	1997	16	3044	Organometallics	HCAPLUS
den Haan, K	1986	15	1726	Organometallics	HCAPLUS
den Haan, K	1986		1726	Organometallics	HCAPLUS
Evans, W	1988	127	575	Inorg Chem	HCAPLUS
Evans, W	1995	134	5927	Inorg Chem	HCAPLUS
Evans, W	11993	12	2618	Organometallics	HCAPLUS
Evans, W	1994	113	1281	Organometallics	HCAPLUS
Ewen, J	1988	110	16255	J Am Chem Soc	HCAPLUS
Ewen, J	1995	l	199	Ziegler Catalysts	HCAPLUS
Giardello, M	1994	116	10212	J Am Chem Soc	HCAPLUS
Giardello, M	1994	116	10212	J Am Chem Soc	HCAPLUS
Heijden, H	1989	•	1255	Organometallics	1
Herrmann, W	1997	16	1813	Organometallics	HCAPLUS
Herrmann, W	1997	16	1813	Organometallics	HCAPLUS
Inoe, N	1993		1	JP 05059077 A2	HCAPLUS
Konemann, M	1997		11155	J Am Chem Soc	1
Kowala, C	1976	B32	1820	Acta Crystallogr	HCAPLUS
Kowala, C	1974		1993	J Chem Soc Chem Comm	HCAPLUS
Lauher, J	1976	98	1729	J Am Chem Soc	HCAPLUS
Lee, L	1994	33	5302	Inorg Chem	HCAPLUS
Lee, M	1998	561	37	J Organomet Chem	HCAPLUS
MoleN	1994	l	1	An Interactive Struc	1
Mosges, G	1992	11	1769	Organometallics	1
Mu, Y	1995	73	2233	Can J Chem	HCAPLUS
Razavi, A	1997	1115	129	,	HCAPLUS
Razavi, A	11995	l	111	Ziegler Catalysts	HCAPLUS
Resconi, L	1998	120	2308	J Am Chem Soc	HCAPLUS
Resconi, L	1996	15	1998	Organometallics	HCAPLUS
Rieger, B	11994	13	1647	Organometallics	HCAPLUS
Rybakova, L	1981	1	1995	Bull Acad Sci USSR D	1
Rybakova, L	1981	1	2415	Izv Akad Nauk SSSR S	HCAPLUS
Schumann, H	1995	496	1233	J Organomet Chem	HCAPLUS
Sharma, R		64	506	•	HCAPLUS
Sheldrick, G	1993	l	1	SHELXL: Program for	1

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Sigalov, A
                        |1983 |
                                     1833
                                            |Bull Acad Sci USSR D|
                                            |Izv Akad Nauk SSSR S|HCAPLUS
Sigalov, A
                        |1983 |
                                     1918
Stern, D
                        |1990 |112
                                     19558
                                            | J Am Chem Soc
                                                                   IHCAPLUS
Tilley, T
                        |1984 |23
                                     12271
                                            |Inorg Chem
                                                                   | HCAPLUS
Westernhausen, M
                        |1995 |621
                                     1837
                                            | Z Anorg Alla Chem
Yoder, J
                        |1998 |17
                                     14946
                                            |Organometallics
                                                                   IHCAPLUS
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L70 ANSWER 14 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

AN 1998:421892 HCAPLUS

DN 129:54646

TI Preliminary investigations on polymerization catalysts composed of lanthanocene and methyl aluminoxane

AU Cui, Li Qiang; Ba, Xiao Wei; Teng, Hong Xiang; Ying, Lai Qiang; Li, Ke Chang; Jin, Ying Tai

CS Changchun Institute Applied Chemistry, Chinese Academy Sciences, Changchun, 130022, Peop. Rep. China

SO Polymer Bulletin (Berlin) (1998), 40(6), 729-734 CODEN: POBUDR; ISSN: 0170-0839

PB Springer-Verlag

DT Journal

LA English

ΙI

AB The polymerization of butadiene (Bd), isoprene (Ip), and styrene (St) was examined using the 6 catalyst systems composed of lanthanocene, (C5H9Cp)2NdCl (I), (C5H9Cp)2SmCl (II), (MeCp)2SmOAr (III), (Ind)2NdCl, Me2Si(Ind)2NdCl (IV), and (Flu)2NdCl, and Me aluminoxane (MAO) resp. All of them can be used to form polyisoprene with mol. wts. of 1000-10,000 and cis-1,4-unit contents of 41-47%. I, II, and III of them can be also used to form polybutadiene with mol. wts. of 10,000-20,000 and cis-1,4-unit contents of 62-78%. In addition, the catalysts from II to IV are still active for St polymerization and

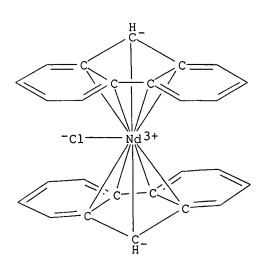
of them gives a syndio-rich random polystyrene. It is noteworthy that II and III are active for homopolymn. of Bd, Ip, and St in the same polymerization condition.

IT 188405-00-7

RL: CAT (Catalyst use); USES (Uses)
(properties of lanthanocene catalysts for polymerization of butadiene or isoprene or styrene)

RN 188405-00-7 HCAPLUS

CN Neodymium, chlorobis[(4a,4b,8a,9,9a-η)-9H-fluoren-9-yl]- (9CI) (CA INDEX NAME)



L70 ANSWER 15 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

AN 1997:219311 HCAPLUS

DN 126:225597

TI Study on polymerization catalysts composed of lanthanocene and methylaluminoxane

AU Cui, Liqiang; Jin, Yingtai; Sun, Junquan; Li, Kechang; Ba, Xiaowei; Teng, Hongxiang

CS Changchun Institute of Applied Chemistry, Chinese Academy of Sciences, 130022, Peop. Rep. China

SO Hecheng Xiangjiao Gongye (1997), 20(2), 79-82 CODEN: HXGOEA; ISSN: 1000-1255

PB Lanzhou Huaxue Gongye Gongsi Huagong Yanjiuyuan

DT Journal

LA Chinese

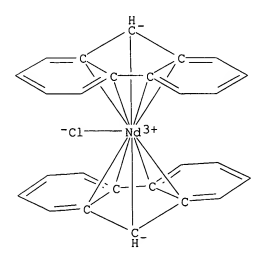
AB Polymns. of butadiene, isoprene, and styrene were carried out using six catalyst systems composed of lanthanocene, (C5H9Cp)2NdCl (I), (C5H9Cp)2SmCl (II), (MeCp)2SmOAr' (III, OAr' is p-methyl-2,6-di-tert-butylphenoxy) (Ind)2NdCl, Me2Si(Ind)2NdCl and (Flu)2NdCl, with methylaluminoxane (MAO) resp. All of catalysts could be used to obtain polyisoprene with relative mol. weight of 1000-10,000 and cis-1,4-unit content of 41-47%. I, II, and III combined with MAO could be used to obtain polybutadiene with relative mol. weight of 10,000-20,000 and cis-1,4-unit content of 62-78%. Polymerization of styrene with II- or III-MAO systems gave polystyrene with high syndiotacticity content.

IT 188405-00-7

RL: CAT (Catalyst use); USES (Uses)
(lanthanocene-methylaluminoxane catalysts for polymerization of butadiene and isoprene and styrene)

RN 188405-00-7 HCAPLUS

CN Neodymium, chlorobis[(4a, 4b, 8a, 9, 9a-η)-9H-fluoren-9-yl]- (9CI) (CA INDEX NAME)



L70 ANSWER 16 OF 16 HCAPLUS COPYRIGHT 2006 ACS on STN

AN 1995:999796 HCAPLUS

DN 124:88161

TI Organometallic catalysts containing cyclopentadienyl group-containing fluorene derivatives and their use for polymerization of vinyl monomers

IN Yasuda, Hajime; Ihara, Eiji; Tokimitsu, Tooru

PA Mitsubishi Rayon Co, Japan

SO Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE			
PΙ	JP 07258319	A2	19951009	JP 1994-52217	19940323 <			
PRAI	JP 1994-52217		19940323	<				
os	MARPAT 124:88161							

GI

$$R^{2}$$
 R^{3}
 R^{4}
 R^{8}
 R^{7}
 R^{10}
 R^{8}
 R^{7}
 R^{10}

Compds. I [M = Sc, Y, lanthanide; R1-8 = H, (Si-containing) C1-5 hydrocarbyl; R9 = CR11R12, SiR11R12; R11, R12 = H, C1-3 alkyl, alkylsilyl; R10 = H, C1-10 hydrocarbyl, alkylsilyl; X = mol. of solvents; m = 0-3; n = 1-3] are prepared and used as catalysts for the polymerization of vinyl monomers.

Ι

Polymerization

of 9.36 mmol Me methacrylate at 0° for 3 h in the presence of 0.10 mmol I [M = Y; R1, R3-6, R8 = H; R2, R7 = tert-Bu; R9 = CMe2; n = 1; R10 = (Me3Si)2CH; m = 0] [prepared from YCl3, 2,7-di-tert-butyl-9-(2-cyclopentadienyl-2-propyl)fluorene Li salt, and (Me3Si)2CHLi] gave a polymer with no-average mol. weight 512,000, mol. weight polydispersity 1.66, and good stereospecificity.

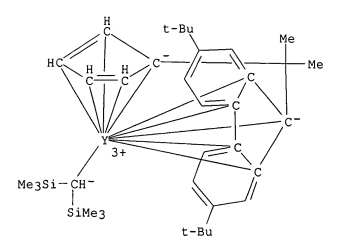
IT 172787-45-0P

RL: CAT (Catalyst use); IMF (Industrial manufacture); PREP (Preparation); USES (Uses)

(catalyst; preparation and use for polymerization of vinyl polymers)

RN 172787-45-0 HCAPLUS

CN Yttrium, [n10-[2,7-bis(1,1-dimethylethyl)-9H-fluoren-9-ylidene](1-methylethylidene)-2,4-cyclopentadien-1-ylidene][bis(trimethylsilyl)methyl)-(9CI) (CA INDEX NAME)



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L71 ANSWER 1 OF 4 HCAPLUS COPYRIGHT 2006 ACS on STN
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AN 2005:1067539 HCAPLUS

DN 143:347612

TI Preparation of styrene homopolymers and styrene-ethylene copolymers using lanthanide metallocene catalyst

IN Carpentier, Jean-Francois; Kirillov, Evgueni;
Razavi, Abbas

PA Total Petrochemicals Research Feluy, Belg.; Centre National de la Recherche Scientifique CNRS

SO Eur. Pat. Appl., 26 pp.

CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 1

FAN.	CNT 1																	
	PATENT NO.			KIND DATE				APPLICATION NO.						DATE				
						-												
ΡI	EP 1582536			A1 20051005			EP 2004-290847					20040331						
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	PL,	SK	
	WO 200	WO 2005095470			A1 20051013			WO 2005-EP51369					20050324					
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	ΒZ,	CA,	CH,	
		CN,	co,	CR,	CU,	CZ,	DE,	DK,	DM,	DŻ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	KP,	KR,	ΚZ,	LC,	
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NA,	NI,	
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	
		SY,	ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	zw
	RV	: BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	
		ΑŻ,	BY,	KG,	ΚZ,	MD,	RU,	ТJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	
		EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	IS,	IT,	LT,	LU,	MC,	NL,	PL,	PT,	
		RO,	SE,	SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	
		MR,	ΝE,	SN,	TD,	TG												
PRAI	EP 200	4-290	847		Α		2004	0331										
os	MARPAT	143:	3476	12														

AB A catalyst system for the homo- or co-polymerization of styrene comprises a metallocene catalyst component of the general formula (Flu-R"-Cp)M(η 3-C3R'5)(ether)n, wherein Cp is a cyclopentadienyl, substituted or unsubstituted, Flu is a fluorenyl, substituted or unsubstituted, M is a

metal Group III of the Periodic Table, ether is a donor solvent mol., R" is a structural bridge between Cp and Flu (9-position) imparting stereorigidity to the component, each R' is the same or different and is hydrogen or an hydrocarbyl having from 1 to 20 carbon atoms and n is 0, 1 or 2. Styrene-ethylene copolymer having a high styrene content was also prepared with the above catalyst.

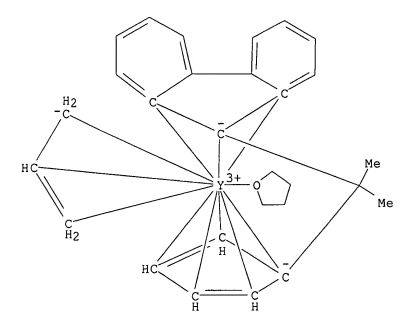
IT 786711-17-9P 866006-06-6P

RL: CAT (Catalyst use); PRP (Properties); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

(preparation of styrene homopolymers and styrene-ethylene copolymers using lanthanide metallocene catalyst)

RN 786711-17-9 HCAPLUS

CN Yttrium, [η8-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene](η3-2-propenyl)(tetrahydrofuran)- (9CI) (CA INDEX NAME)



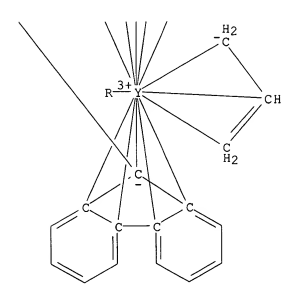
RN 866006-06-6 HCAPLUS

CN Yttrium, [η10-[3-(1,1-dimethylethyl)-2,4-cyclopentadien-1-ylidene](1methylethylidene)-9H-fluoren-9-ylidene](η3-2propenyl)(tetrahydrofuran)- (9CI) (CA INDEX NAME)

PAGE 1-A

$$\begin{array}{c} \text{Me} \\ \text{Me} \\ \text{C} \\$$

PAGE 2-A



PAGE 3-A



IT 866006-04-4P

RL: CAT (Catalyst use); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

(preparation of styrene homopolymers and styrene-ethylene copolymers using lanthanide metallocene catalyst)

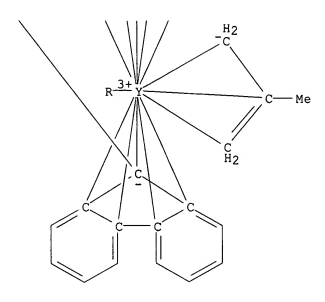
RN 866006-04-4 HCAPLUS

CN Yttrium, $[\eta 10-2, 4-\text{cyclopentadien}-1-\text{ylidene}(1-\text{methylethylidene})-9H-fluoren-9-ylidene][(1,2,3-<math>\eta$)-2-methyl-2-propenyl](tetrahydrofuran)-(9CI) (CA INDEX NAME)

PAGE 1-A

$$\begin{array}{c} \text{Me} \\ \text{Me} \\ \end{array} \begin{array}{c} -\text{C} \\ \end{array} \begin{array}{c} \text{H} \\ \text{C} \\ \end{array} \begin{array}{c} \text{CH} \\ \end{array}$$

PAGE 2-A



PAGE 3-A

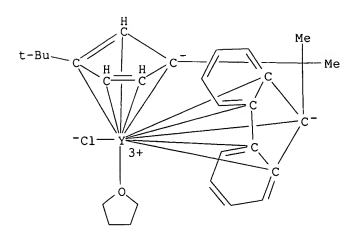


IT 706760-93-2P

RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation of styrene homopolymers and styrene-ethylene copolymers using lanthanide metallocene catalyst)

RN 706760-93-2 HCAPLUS

CN Yttrium, chloro[η 10-[3-(1,1-dimethylethyl)-2,4-cyclopentadien-1-ylidene](1-methylethylidene)-9H-fluoren-9-ylidene](tetrahydrofuran)- (9CI) (CA INDEX NAME)



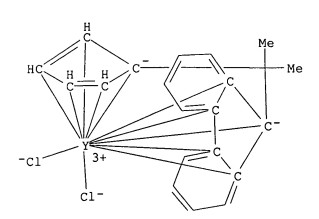
IT 611233-16-0

RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of styrene homopolymers and styrene-ethylene copolymers using lanthanide metallocene catalyst)

RN 611233-16-0 HCAPLUS

CM 1

CRN 611233-15-9 CMF C21 H18 C12 Y CCI CCS



CM 2

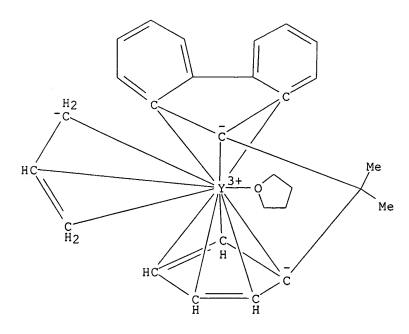
CRN 444121-94-2 CMF C16 H34 Li O4

CCI CCS

RETABLE

Referenced Author (RAU)	(RPY) (RVL) (RPG)	•	Referenced File
Anon	1996 1996	•	PATENT ABSTRACTS OF	
Carpentier	12004	1	WO 2004060942 A	HCAPLUS
Ipsco Inc	1996	1	WO 9607861 A	1
Lu, Z	1994 53	1453	J APPL POLYM SCI	HCAPLUS
Mitsubishi Rayon Co Lt		1	JP 07258319 A	HCAPLUS
Mitsubishi Rayon Co Lt	d 1995	1	JP 7258319 A	
Sernetz	1997 35	1571	JOURNAL OF POLYMER S	HCAPLUS

- L71 ANSWER 2 OF 4 HCAPLUS COPYRIGHT 2006 ACS on STN
- AN 2004:729326 HCAPLUS
- DN 141:395882
- TI Highly Syndiospecific Polymerization of Styrene Catalyzed by Allyl Lanthanide Complexes
- AU Kirillov, Evgueni; Lehmann, Christian W.; Razavi, Abbas; Carpentier, Jean-Francois
- CS Organometalliques et Catalyse, UMR 6509, Institut de Chimie de Rennes, CNRS-Universite de Rennes 1, Rennes, 35042, Fr.
- SO Journal of the American Chemical Society (2004), 126(39), 12240-12241 CODEN: JACSAT; ISSN: 0002-7863
- PB American Chemical Society
- DT Journal
- LA English
- OS CASREACT 141:395882
- AB Allylic complexes of lanthanides bearing a fluorenyl-based ligand are active single-component catalysts for the polymerization of styrene, giving highly syndiotactic polymers (rrrr > 99%) with low to high mol. weight (Mn = 8000-135 000) and narrow polydispersities (Mw/Mn = 1.25-2.1).
- IT 786711-17-9P
 - RL: CAT (Catalyst use); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)
 - (highly syndiospecific polymerization of styrene catalyzed by allyl lanthanide complexes)
- RN 786711-17-9 HCAPLUS
- CN Yttrium, [η 8-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene](η 3-2-propenyl)(tetrahydrofuran)- (9CI) (CA INDEX NAME)



IT **611233-16-0**

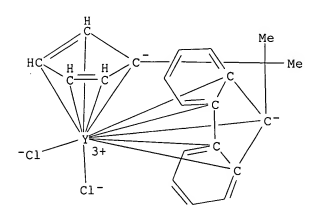
RL: RCT (Reactant); RACT (Reactant or reagent)
 (highly syndiospecific polymerization of styrene catalyzed by allyl lanthanide
 complexes)

RN 611233-16-0 HCAPLUS

CN Lithium(1+), [1,1'-oxybis[ethane]]tris(tetrahydrofuran)-, (T-4)-, dichloro[η 10-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene]yttrate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 611233-15-9 CMF C21 H18 C12 Y CCI CCS



CM 2

CRN 444121-94-2

CMF C16 H34 Li O4 CCI CCS

RE	ΤА	$_{ m BL}$	Ε

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	Year	VOL	PG	Referenced Work	Referenced
(RAU)	(RPY)			(RWK)	File
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- L71 ANSWER 3 OF 4 HCAPLUS COPYRIGHT 2006 ACS on STN
- AN 2004:325442 HCAPLUS
- DN 141:54425
- TI Synthesis, Structure, and Polymerization Activity of Neutral Halide, Alkyl, and Hydrido Yttrium Complexes of Isopropylidene-Bridged Cyclopentadienyl-Fluorenyl Ligands
- AU Kirillov, Evgueni; Lehmann, Christian W.; Razavi, Abbas; Carpentier, Jean-Francois
- CS Organometalliques et Catalyse, UMR 6509, Institut de Chimie de Rennes, CNRS-Universite de Rennes 1, Rennes, 35042, Fr.
- SO Organometallics (2004), 23(11), 2768-2777

CODEN: ORGND7; ISSN: 0276-7333

PB American Chemical Society

DT Journal LA English

OS CASREACT 141:54425

AB Reactions of the anionic complex [(Cp-CMe2-Flu)YCl2]-[Li(ether)4]+ (1) (Cp = C5H4, Flu = 9-C13H8), prepared in situ from YCl3(THF)3.5 and 1 molar equivalent of the dilithium salt [Cp-CMe2-Flu]Li2, with equimolar amts. of RLi give alkyl mono-THF complexes [(Cp-CMe2-Flu)]Y(R)(THF) (R = CH(SiMe3)2, 3; CH2SiMe3, 4) in high yields. The solid-state structure of 3 was established by x-ray diffraction, showing the fluorenyl moiety sym. coordinated to yttrium in an intermediary $\eta 3-\eta 5$ mode. Hydrogenolysis of 3 and 4 with H2 or PhSiH3 gives the hydride $\{[(\mu:\eta 5,\eta 5-Cp-CMe2-Flu)]Y(\mu-H)(THF)\}2$ (5). The solid-state structure of 5 was determined by x-ray diffraction, revealing a dimeric structure with both bridging Cp-CMe2-Flu and hydride ligands (Y-H = 1.99(4)-2.01(4) Å). Complex 5 is the first structurally characterized example of a group 3 metal hydride stabilized by a fluorenyl ligand. Reaction of 1 with PhCH2MqBr gives, instead of a benzyl derivative, the neutral base-free bromo complex { $[(\eta 5, \eta 5-Cp-CMe2-Flu)]Y(\mu-Br)}2$ (6), which shows a dimeric structure in the solid state with chelating Cp-CMe2-Flu and bridging bromide ligands. Introduction of the bulky tert-Bu substituent on the Cp ring of the ligand system enabled the preparation of the neutral chloro complex [(3-tBuCp)-CMe2-Flu]YCl(THF) (7), using a salt elimination between the dilithium salt of the ligand and YCl3(THF)3.5. Reaction of 7 with LiCH(SiMe3)2 gives the alkyl complex {[(3-tBuCp)-CMe2-Flu]Y(CH(SiMe3)2)} (8), which contains no THF mol. in its coordination sphere in contrast to unsubstituted analogs 3 and 4. Preliminary studies of the catalytic activity of these new complexes for ethylene and MMA polymerization are reported.

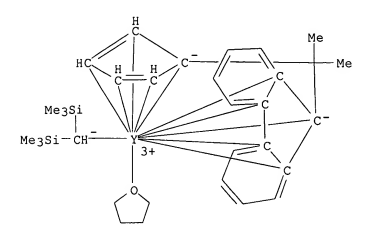
IT 706760-85-2P

RL: CAT (Catalyst use); PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (crystal structure; preparation, structure, and polymerization catalytic activity

of neutral halide, alkyl, and hydrido yttrium complexes of isopropylidene-bridged cyclopentadienyl-fluorenyl ligands)

RN 706760-85-2 HCAPLUS

CN Yttrium, [bis(trimethylsilyl)methyl][\(\eta 10-2\), 4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene](tetrahydrofuran)- (9CI) (CA INDEX NAME)



IT 706760-97-6P 708211-99-8P

of neutral halide, alkyl, and hydrido yttrium complexes of isopropylidene-bridged cyclopentadienyl-fluorenyl ligands)

RN 706760-97-6 HCAPLUS

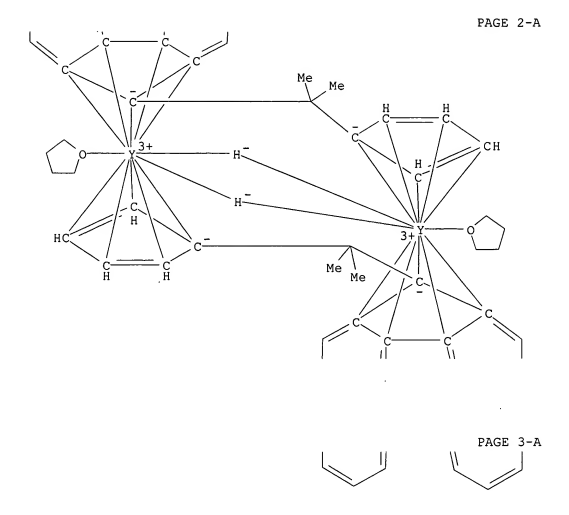
CN Yttrium, bis[μ -[η 5: η 5-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene]]di- μ -hydrobis(tetrahydrofuran)di-, compd. with benzene (2:1) (9CI) (CA INDEX NAME)

CM 1

CRN 706760-89-6 CMF C50 H54 O2 Y2 CCI CCS

PAGE 1-A





CM 2

CRN 71-43-2 CMF C6 H6

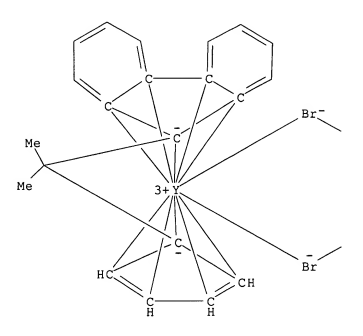


RN 708211-99-8 HCAPLUS
CN Yttrium, di-μ-bromobis[η10-2,4-cyclopentadien-1-ylidene(1 methylethylidene)-9H-fluoren-9-ylidene]di-, stereoisomer, compd. with
 methylbenzene (1:1) (9CI) (CA INDEX NAME)

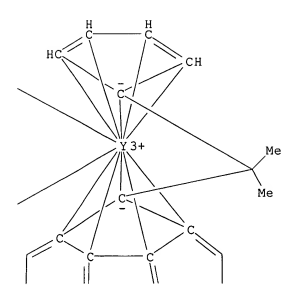
CM 1

CRN 706760-91-0 CMF C42 H36 Br2 Y2 CCI CCS

PAGE 1-A



PAGE 1-B



PAGE 2-B

" "

CM 2

CRN 108-88-3 CMF C7 H8

IT 706760-89-6P 706760-91-0P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (mol. structure; preparation, structure, and polymerization catalytic activity

of
neutral halide, alkyl, and hydrido yttrium complexes of
isopropylidene-bridged cyclopentadienyl-fluorenyl ligands)

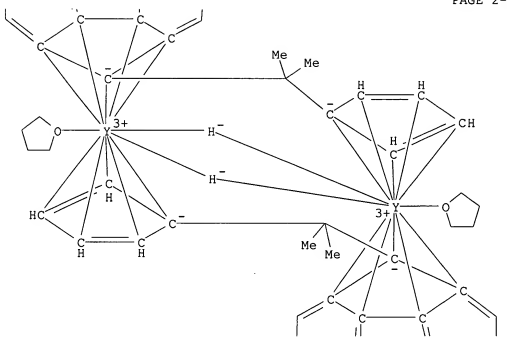
RN 706760-89-6 HCAPLUS

CN Yttrium, bis $[\mu-[\eta 5:\eta 5-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene]]di-<math>\mu$ -hydrobis(tetrahydrofuran)di-(9CI) (CA INDEX NAME)

PAGE 1-A





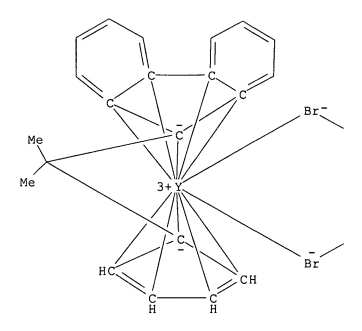




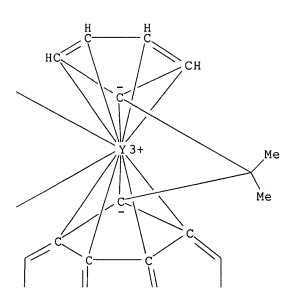
RN 706760-91-0 HCAPLUS

CN Yttrium, di- μ -bromobis[η 10-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene]di-, stereoisomer (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 1-B



PAGE 2-B



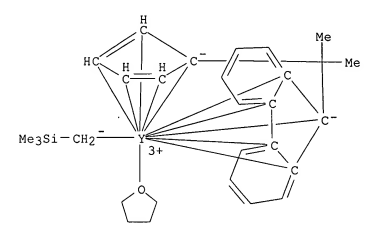
IT 706760-87-4P

RL: CAT (Catalyst use); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (preparation, structure, and polymerization catalytic activity of neutral

halide,
alkyl, and hydrido yttrium complexes of isopropylidene-bridged
cyclopentadienyl-fluorenyl ligands)

RN 706760-87-4 HCAPLUS

CN Yttrium, [n10-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9H-fluoren-9-ylidene)(tetrahydrofuran)[(trimethylsilyl)methyl]- (9CI) (CA INDEX NAME)



IT 611233-16-0

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation, structure, and polymerization catalytic activity of neutral halide,

alkyl, and hydrido yttrium complexes of isopropylidene-bridged cyclopentadienyl-fluorenyl ligands)

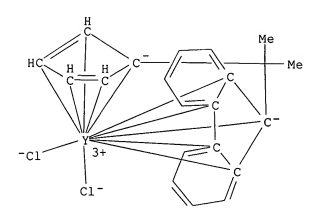
RN 611233-16-0 HCAPLUS

CN Lithium(1+), [1,1'-oxybis[ethane]]tris(tetrahydrofuran)-, (T-4)-,
dichloro[η10-2,4-cyclopentadien-1-ylidene(1-methylethylidene)-9Hfluoren-9-ylidene]yttrate(1-) (9CI) (CA INDEX NAME)

CM 1

CRN 611233-15-9 CMF C21 H18 C12 Y

CCI CCS



CM 2

CRN 444121-94-2 CMF C16 H34 Li O4 CCI CCS

IT 706760-83-0P 706760-93-2P 706760-95-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation, structure, and polymerization catalytic activity of neutral halide,

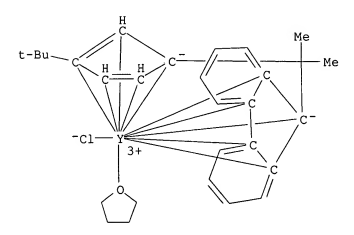
alkyl, and hydrido yttrium complexes of isopropylidene-bridged cyclopentadienyl-fluorenyl ligands)

RN 706760-83-0 HCAPLUS

CN Yttrium, bis[$(1,2,3,4,5-\eta)-1-[1-(9H-fluoren-9-yl)-1-methylethyl]-2,4-cyclopentadien-1-yl]bis(tetrahydrofuran)[(trimethylsilyl)methyl]- (9CI) (CA INDEX NAME)$

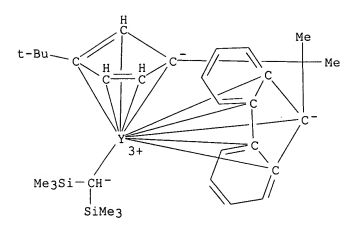
RN 706760-93-2 HCAPLUS

Yttrium, chloro[n10-[3-(1,1-dimethylethyl)-2,4-cyclopentadien-1-ylidene](1-methylethylidene)-9H-fluoren-9-ylidene](tetrahydrofuran)- (9CI)
(CA INDEX NAME)



RN 706760-95-4 HCAPLUS

CN Yttrium, [bis(trimethylsilyl)methyl][\(\eta 10 - [3 - (1, 1 - dimethylethyl) - 2, 4 - cyclopentadien - 1 - ylidene](1 - methylethylidene) - 9H - fluoren - 9 - ylidene] - (9CI) (CA INDEX NAME)



RAU RPY RVL RPG RWK File	RETABLE					
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ΑN
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TΙ
     [\eta 5: \eta 1-(3, 6-tBu2Flu) SiMe2NtBu] Y (\eta 1-NC5H6) (py) 2: a
     1,4-hydride-addition product to pyridine that provides evidence for the
     first fluorenyl(hydrido)metal (Group 3) complex
ΑU
     Kirillov, Evgueni; Lehmann, Christian W.; Razavi, Abbas
     ; Carpentier, Jean-Francois
     Organometalliques et Catalyse, Institut de Chimie de Rennes, UMR 6509
CS
     CNRS-Universite de Rennes 1, Rennes, 35042, Fr.
     European Journal of Inorganic Chemistry (2004), (5), 943-945
SO
     CODEN: EJICFO; ISSN: 1434-1948
PB
     Wiley-VCH Verlag GmbH & Co. KGaA
DT
     Journal
LA
     English
OS
     CASREACT 141:157248
     Reaction of pyridine with the constrained geometry
AΒ
     fluorenyl(hydrido)yttrium complex {[(3,6-tBu2Flu)SiMe2NtBu]Y(H)(THF)}2
     selectively gives the 1,4-addition product [\eta 5:\eta 1-(3,6-
     tBu2Flu)SiMe2NtBu]Y(η1-NC5H6)(py)2, which was characterized by
     single-crystal x-ray diffraction and 1H and 13C NMR spectroscopy.
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RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation and crystal structure of 1,4-hydride-addition product of pyridine with yttrium hydrido fluorenyl complex)

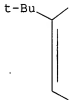
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CN Yttrium, bis[1-[(4a,4b,8a,9,9a- η)-3,6-bis(1,1-dimethylethyl)-9H-fluoren-9-yl]-N-(1,1-dimethylethyl)-1,1-dimethylsilanaminato(2-)- κ N]di- μ -hydrobis(tetrahydrofuran)di- (9CI) (CA INDEX NAME)

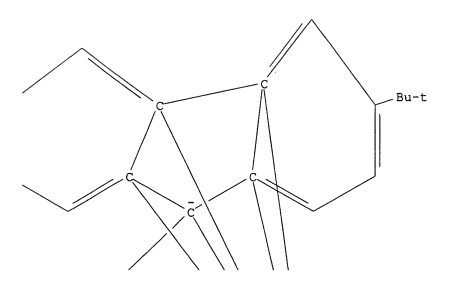
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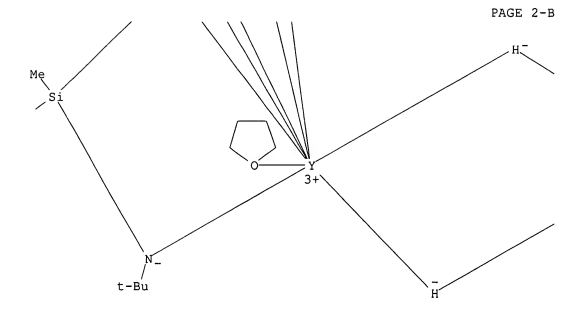


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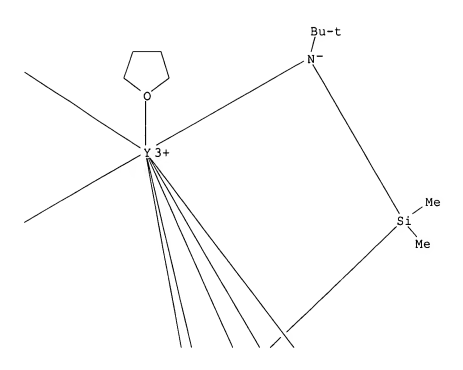


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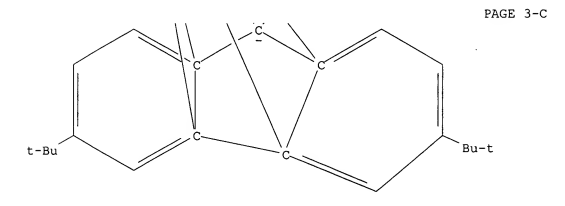


PAGE 2-C



PAGE 3-A

| HCAPLUS



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jan delaval - 4 october 2006

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